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(54) Title: PEPTIDE NUCLEIC ACIDS

(57) Abstract

A novel class of compounds, known as peptide nucleic acids, bind complementary ssDNA and RNA strands more strongly than a corresponding DNA. The peptide nucleic acids generally comprise ligands such as naturally occurring DNA bases attached to a peptide backbone through a suitable linker.

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PEPTIDE NUCLEIC ACIDS

RELATED APPLICATIONS

This application is a continuation-in-part of the following Danish Patent Applications: No. 986/91, filed May 24, 1991, No. 987/91, filed May 24, 1991, and No. 510/92, filed April 15, 1992. The entire disclosure of each application is incorporated herein by reference.

10 FIELD OF THE INVENTION

This invention is directed to compounds that are not polynucleotides yet which bind to complementary DNA and RNA strands more strongly the corresponding DNA. In particular, the invention concerns compounds wherein naturally-occurring nucleobases or other nucleobase-binding moieties ar covalently bound to a polyamide backbone.

BACKGROUND OF THE INVENTION

Oligodeoxyribonucleotides as long as 100 base pairs 20 (bp) are routinely synthesized by solid phase methods using commercially available, fully automatic synthesis machines. The chemical synthesis of oligoribonucleotides, however, is far less routine. Oligoribonucleotides also are much less stable than oligodeoxyribonucleotides, a fact which has contributed to the more prevalent use of oligodeoxyribonucl otides in medical and biological research directed to, for xample, gene therapy or the regulation of transcription r translation.

The function of a g ne starts by transcripti n f its information to a messenger RNA (mRNA) which, by interaction with th ribosomal complex, directs the synthesis of a protein coded for by its sequence. The synthetic process is known as 5 translation. Translation requires the presence of various cofactors and building blocks, the amino acids, and their transfer RNAs (tRNA), all of which are present in normal cells.

Transcription initiation requires specific recognition
of a promoter DNA sequence by the RNA-synthesizing enzyme, RNA
polymerase. In many cases in prokaryotic cells, and probably
in all cases in eukaryotic cells, this recognition is preceded
by sequence-specific binding of a protein transcription factor
to the promoter. Other proteins which bind to the promoter,
but whose binding prohibits action of RNA polymerase, are
known as repressors. Thus, gene activation typically is
regulated positively by transcription factors and negatively
by repressors.

Most conventional drugs function by interaction with 20 and modulation of one or more targeted endogenous proteins, Such drugs, however, typically are not e.g., enzymes. specific for targeted proteins but interact with other proteins as well. Thus, a relatively large dose of drug must be used to effectively modulate a targeted protein. Typical 25 daily doses of drugs are from 10°5-10°1 millimoles per kilogram of body weight or 10⁻³-10 millimoles for a 100 kilogram If this modulation instead could be effected by interaction with and inactivation of mRNA, a dramatic reduction in the necessary amount of drug necessary could 30 likely be achieved, along with a corresponding reduction in side effects. Further reductions could be effected if such interaction could be rendered site- specific. Given that a functioning gene continually produces mRNA, it would thus be even more advantageous if gene transcription could be arrested 35 in its entirety.

Oligodeoxynucleotides offer such pportunities. For example, synthetic oligodeoxynucleotides could be used as

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antisense probes to block and ev ntually lead to the breakdown of mRNA. Thus, synthetic DNA c uld suppress translation in vivo. It also may be possible to modulate the genome f an animal by, for example, triple helix formation using oligonucleotides or other DNA recognizing agents. However, there are a number of drawbacks associated with triple helix formation. For example, it can only be used for homopurin sequences and it requires unphysiologically high ionic strength and low pH.

10 Furthermore, unmodified oligonucleotides ar unpractical both in the antisense approach and in the triple helix approach because they have short in vivo half-lives, they are difficult to prepare in more than milligram quantities and, thus, are prohibitively costly, and they are 15 poor cell membrane penetrators.

These problems have resulted in an extensive search for improvements and alternatives. For example, the problems arising in connection with double-stranded DNA (dsDNA) recognition through triple helix formation have been diminished by a clever "switch back" chemical linking whereby a sequence of polypurine on one strand is recognized, and by "switching back", a homopurine sequence on the other strand can be recognized. See, e.g., McCurdy, Moulds, and Froehler, Nucleosides, in press. Also, good helix formation has been obtained by using artificial bases, thereby improving binding conditions with regard to ionic strength and pH.

In order to improve half life as well as membrane penetration, a large number of variations in polynucleotide backbones has been undertaken, although so far not with 30 desired results. These variations include the use f methylphosphonates, monothiophosphates, dithiophosphates, phosphoramidates, phosphate esters, bridged phosphoroamidates, bridged phosphorothioates, bridged methylenephosphonates, dephospho internucleotide analogs with siloxane bridges, carbonate bridges, carboxymethyl ester bridges, acetamid bridges, carbamat bridges, thi ther, sulfoxy, sulf no

bridges, various "plastic" DNAs, α -anomeric bridges, and borane derivatives.

International patent applicati n WO 86/05518 broadly claims a polymeric composition effective to bind to a single-5 stranded polynucleotide containing a target sequence of bases. The composition is said to comprise non-homopolymeric, substantially stereoregular polymer molecules of the form:

$$R_1$$
 R_2 R_3 R_n
 $B - B - B - \dots$ B_n

10 where:

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- (a) R₁-R_n are recognition moieties selected from purine, purine-like, pyrimidine, and pyrimidine like heterocycles effective to bind by Watson/Crick pairing to corresponding, in-sequence bases in the target sequence;
- (b) n is such that the total number of Watson/Crick hydrogen bonds formed between a polymer molecule and target sequence is at least about 15;
- (c) B B are backbone moieties joined predominantly by chemically stable, substantially uncharged, predominantly achiral linkages;
 - (d) the backbone moiety length ranges from 5 to 7 atoms if the backbone moieties have a cyclic structure, and ranges from 4 to 6 atoms if the backbone moieties have an acyclic structure; and
 - (e) the backbone moieties support the recognition moieties at position which allow Watson/Crick base pairing between the recognition moieties and the corresponding, in-sequence bases of the target sequence.

According to WO 86/05518, the recognition moieties are various natural nucleobases and nucleobase-analogs and the backbone moieties are either cyclic backbone moieties comprising furan or morpholine rings or acyclic backbone moieties of the 35 following forms:

where E is -CO- or -SO₂-. The specification of the application provides general descriptions for the synthesis of subunits, for backbone coupling reactions, and for polymer assembly strategies. However, the specification provides n example wherein a claimed compound or structure is actually prepared. Although WO 86/05518 indicates that the claim d polymer compositions can bind target sequences and, as a result, have possible diagnostic and therapeutic applications, the application contains no data relating to the binding affinity of a claimed polymer.

International patent application WO 86/05519 claims diagnostic reagents and systems that comprise polymers 40 described in WO 86/05518, but attached to a solid support. WO 86/05519 also provides no examples concerning actually preparation of a claimed diagnostic reagent, much less data showing the diagnostic efficiency of such a reagent.

International patent application WO 89/12060 claims 45 various building blocks for synth sizing oligonucleotide analogs, as well as lig nucleotide analogs formed by joining

such building blocks in a defined sequence. The building blocks may be either "rigid" (containing a ring) or "flexible" (lacking a ring). In both cases the building blocks contain a hydroxy group and a mercapto group, through which the 5 building blocks are said to join to form oligonucleotide analogs. The linking moiety in the oligonucleotide analogs is selected from the group consisting of -sulfide (-S-), sulfoxide (-SO-), and sulfone (-SO₂-). WO 89/12060 provides a general description concerning synthesis of the building 10 blocks and coupling reactions for the synthesis of oligonucleotide analogs, along with experimental examples describing the preparation of building blocks. However, the application provides no examples directed to the preparation of a claimed oligonucleotide analog and no data confirming the 15 specific binding of an oligonucleotide analog to a target oligonucleotide.

Furthermore, oligonucleotides or their derivatives have been linked to intercalators in order to improve binding, to polylysine or other basic groups in order to improve binding 20 both to double-stranded and single-stranded DNA, and to peptides in order to improve membrane penetration. However, such linking has not resulted in satisfactory binding for either double-stranded or single-stranded DNA. Other problems which resulted from, for example, methylphosphonates and 25 monothiophosphates were the occurrence of chirality, insufficient synthetic yield or difficulties in performing solid phase assisted syntheses.

In most cases only a few of these modifications could be used. Even then, only short sequences -- often only dimers -- or monomers could be generated. Furthermore, the oligomers actually produced have rarely been shown to bind to DNA or RNA or have not been examined biologically.

The great majority of these backbone modifications led to decreased stability for hybrids formed between the modified oligonucleotide and its complementary native oligonucleotide, as assayed by measuring T_a values. Consequently, it is generally understood in the art that backbone modifications

destabilize such hybrids, i. ., result in low r T_n valu s, and should be kept to a minimum.

OBJECTS OF THE INVENTION

It is one object of the present invention to provid compounds that bind ssDNA and RNA strands to form stable hybrids therewith.

It is a further object of the invention to provid compounds that bind ssDNA and RNA strands more strongly th 10 corresponding DNA.

It is another object to provide compounds wherein naturally-occurring nucleobases or other nucleobase-binding moieties are covalently bound to a peptide backbone.

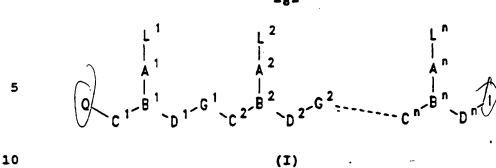
It is yet another object to provide compounds oth r 15 than RNA that can bind one strand of a double-strand d polynucleotide, thereby displacing the other strand.

It is still another object to provide therapeutic and prophylactic methods that employ such compounds.

20 SUMMARY OF THE INVENTION

The present invention provides a novel class of compounds, known as peptide nucleic acids (PNAs), that bind complementary ssDNA and RNA strands more strongly than a corresponding DNA. The compounds of the invention generally comprise ligands linked to a peptide backbone via an aza nitrogen. Representative ligands include either the four main naturally occurring DNA bases (i.e., thymine, cytosine, adenine or guanine) or other naturally occurring nucleobases (e.g., inosine, uracil, 5-methylcytosine or thiouracil) or artificial bases (e.g., bromothymine, azaadenines or azaguanines, etc.) attached to a peptide backbone through a suitable linker.

In certain preferred embodiments, the peptide nucleic acids of the invention have the general formula (I):



wherein:

n is at least 2,

each of L¹-L¹ is independently selected from the group consisting of hydrogen, hydroxy, (C₁-C₄)alkanoyl, naturally occurring nucleobases, non-naturally occurring nucleobases, aromatic moieties, DNA intercalators, nucleobase-binding groups, heterocyclic moieties, and reporter ligands, at least one of L¹-L¹ being a naturally occurring nucleobase, a non-naturally occurring nucleobase, a DNA intercalator, or a nucleobase-binding group;

each of A¹-Aⁿ is a single bond, a methylene group or a group of formula (IIa) or (IIb):

25 $\begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}_p \begin{bmatrix} R^1 \\ R^2 \end{bmatrix}_q \quad \text{or} \quad \begin{bmatrix} R^1 \\ C \\ R^2 \end{bmatrix}_r \begin{bmatrix} R^1 \\ R^2 \end{bmatrix}_s$ (IIa) (IIb)

where:

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X is O, S, Se, NR^3 , CH_2 or $C(CH_3)_2$; Y is a single bond, O, S or NR^4 ;

each of p and q is zero or an integer from 1 to 5, the sum p+q being not more than 10;

each of r and s is zero or an integer from 1 to 5, the sum r+s being not more than 10;

each R^1 and R^2 is independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl which may be hydroxy- or alkoxy- r alkylthi -substitut d, hydroxy, alkoxy, alkylthio, amino and halogen; and

each R^3 and R^4 is ind pendently s 1 ct d from the group consisting f hydrog n, (C_1-C_4) alkyl, hydr xy- r alkoxy- or alkylthi-substitut d (C_1-C_4) alkyl, hydr xy, alkylthio and amino;

each of B¹-Bⁿ is N or R³N², where R³ is as defined above;
each of C¹-Cⁿ is CR⁶R⌉, CHR⁶CHR⌉ or CR⁶R⌉CH₂, where R⁶ is
hydrogen and R⌉ is selected from the group consisting of the
side chains of naturally occurring alpha amino acids, or R⁶
and R⌉ are independently selected from the group consisting of
hydrogen, (C₂-C₆)alkyl, aryl, aralkyl, heteroaryl, hydroxy,
(C₁-C₆)alkoxy, (C₁-C₆)alkylthio, NRȝR⁶ and SRȝ, where Rȝ and R⁶
are as defined above, and Rȝ is hydrogen, (C₁-C₆)alkyl,
hydroxy-, alkoxy-, or alkylthio- substituted (C₁-C₆)alkyl, or
R⁶ and R⌉ taken together complete an alicyclic or heterocyclic
system;

each of D^1-D^n is CR^6R^7 , $CH_2CR^6R^7$ or CHR^6CHR^7 , where R^6 and R^7 are as defined above;

each of G^1-G^{n-1} is $-NR^3CO-$, $-NR^3CS-$, $-NR^3SO-$ or $-NR^3SO_2-$, Y in either orientation, where R^3 is as defined above;

Q is -CO₂H, -CONR'R'', -SO₃H or -SO₂NR'R'' or an activated derivative of -CO₂H or -SO₃H; and

I is -NHR'''R'''' or -NR'''C(0)R'''', where R', R",
R''' and R'''' are independently selected from the group
consisting of hydrogen, alkyl, amino protecting groups,
reporter ligands, intercalators, chelators, peptides,
proteins, carbohydrates, lipids, steroids, oligonucleotides
and soluble and non-soluble polymers.

The peptide nucleic acids of the invention differ fr m those disclosed in WO 86/05518 in that their recogniti n moieties are attached to an aza nitrogen atom in the backbon, rather than to an amide nitrogen atom, a hydrazine moiety r a carbon atom in the backbone.

Preferred peptide nucleic acids have general formula (III):

$$\begin{array}{c|c}
R^{n} & \downarrow & \downarrow \\
CH_{2})_{k} & \downarrow & \downarrow \\
CH_{2})_{k} & \downarrow & \downarrow \\
R^{7} &$$

(III)

wherein:

each L is independently selected from the group consisting of hydrogen, phenyl, heterocyclic moieties, naturally occurring nucleobases, and non-naturally occurring nucleobases;

each R⁷ is independently selected from the group 10 consisting of hydrogen and the side chains of naturally occurring alpha amino acids;

n is an integer from 1 to 60;

each of k, l and m is independently zero or an integer
from 1 to 5;

R^h is OH, NH₂ or -NHLysNH₂; and Rⁱ is H or COCH₁.

Particularly preferred are compounds having formula (III) wherein each L is independently selected from the group consisting of the nucleobases thymine (T), adenine (A), 20 cytosine (C), guanine (G) and uracil (U), k and m are zero or 1, and n is an integer from 1 to 30, in particular from 4 to 20. An example of such a compound is provided in Figure 1, which shows the structural similarity between such compounds and single-stranded DNA.

25 The peptide nucleic acids of the invention are synthesized by adaptation of standard peptide synthesis procedures, either in solution or on a solid phase. The synthons used are specially designed m nomer amino acids or their activat d derivatives, protected by standard protecting

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groups. The ligonucleotid analogs als can be synthesized by using the cresponding diacids and diamines.

Thus, the now less moments and large to the invention are selected from the group consisting of amino acids, diacids and diamines having general formulae:

15 wherein L, A, B, C and D are as defined above, except that any amino groups therein may be protected by amino protecting groups; E is COOH, CSOH, SOOH, SO₂OH or an activat d derivative thereof; and F is NHR³ or NPgR³, where R³ is as defined above and Pg is an amino protecting group.

20 Preferred monomer synthons according to the inventi n are amino acids having formula (VII):

(VII)

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or amino-protected and/or acid terminal activated derivatives thereof, wherein L is selected from the group consisting of hydrogen, phenyl, heterocyclic moieties, naturally occurring nucleobases, non-naturally occurring nucleobases, and protected derivatives thereof; and R' is independently selected from the group consisting of hydrogen and the side chains of naturally occurring alpha amino acids. Especially preferred are such synthons having formula (VII) wherein R' is hydrogen and L is selected from the group consisting of the nucl obas s thymin (T), adenine (A), cytosine (C), guanin (G) and uracil (U) and protect d derivatives thereof.

Unexpectedly, these compounds also are able to recognize duplex DNA by displacing ne strand, thereby presumably generating a double helix with the ther one. Such recognition can take place to dsDNA sequences 5-60 base pairs 5 long. Sequences between 10 and 20 bases are of interest since this is the range within which unique DNA sequences of prokaryotes and eukaryotes are found. Reagents which recognize 17-18 bases are of particular interest since this is the length of unique sequences in the human genome. The 10 compounds of the invention also should be able to form triple helices with dsDNA.

Whereas the improved binding of the compounds of the invention should render them efficient as antisense agents, it is expected that an extended range of related reagents may 15 cause strand displacement, now that this surprising and unexpected new behavior of dsDNA has been discovered.

Thus, in one aspect, the present invention provides methods for inhibiting the expression of particular genes in the cells of an organism, comprising administering to said organism a reagent as defined above which binds specifically to sequences of said genes.

Further, the invention provides methods for inhibiting transcription and/or replication of particular genes or for inducing degradation of particular regions of double stranded 25 DNA in cells of an organism by administering to said organism a reagent as defined above.

Still further, the invention provides methods for killing cells or virus by contacting said cells or virus with a reagent as defined above which binds specifically to 30 sequences of the genome of said cells or virus.

BRIEF DESCRIPTION OF THE DRAWINGS

The numerous obj cts and advantag s f th pres nt invention may be better underst od by th s skilled in the art by reference to the accompanying figures, in which:

Figure 1 shows a naturally occurring deoxyribooligonucleotide (A) and a peptide nucleic acid (PNA) of the invention (B).

Figure 2 provides examples of naturally occurring and non-naturally occurring nucleobases for DNA recognition and 10 reporter groups.

Figure 3 provides a schematic illustration of (a) photocleavage by Acr¹-(Taeg)₁₀-Lys-NH₂ (Acr-T10-LysNH₂); (b) photofootprint by the diazo-linked acridine of Acr¹-(Taeg)₁₀-Lys-NH₂ and preferred KMnO₄-cleavage; and (c) S₁-nuclease 15 enhanced cleavage and (d) micrococcus nuclease cleavage f Acr¹-(Taeg)₁₀-Lys-NH₂ binding site.

Figure 4 provides examples of PNA monomer synthons of the invention.

Figure 5 shows the Acr¹ ligand and a PNA, Acr^{1} -(Taeg)₁₀-20 Lys-NH₂.

Figure 6 provides a general scheme for the preparati n of monomer synthons.

Figure 7 provides a general scheme for the preparation of the Acr¹ ligand.

25 Figure 8 provides a general scheme for solid-phase PNA synthesis illustrating the preparation of linear unprotected PNA amides.

Figure 9 shows analytical HPLC chromatograms of: (A) crude H-[Taeg]₁₅-NH₂ after HF-cleavage (before lyophilization);

- 30 (B) crude Acr¹-[Taeg]₁₅-NH₂ after HF-cleavage (before lyophilization); and (C) purified Acr¹-[Taeg]₁₅-NH₂. Buffer A, 5t CH₃CN/95t H₂O/0.0445t TFA; buffer B, 60t CH₃CN/40t H₂O/0.0390t TFA; linear gradient, 0-100t of B in 30 min; fl w rate, 1.2 ml/min; column, Vydac C₁₈ (5 μm, 0.46 x 25 cm).
- Figure 10 shows analytical HPLC chromatograms of: (A) purified H-[Ta g]₁₀-Lys-NH₂ and (B) purifi d H-[Taeg]₅-Caeg[Taeg]₄-Lys-NH₂ mploying the same conditi ns as in Figure 9.

Figures 11a and 11b show binding of AcrT10-Lys to dA_{10} . $5'-^{32}P$ -labeled oligonucleotide (1) $(5'-GATCCA_{10}G)$ was incubated in the absence or presence of Acr-T10-LysNH₂ and in the absence or presence of oligonucleotide (2) $(5'-GATCCT_{10}G)$ and the samples were analyzed by polyacrylamide gel electrophoresis (PAGE) and autoradiography under "native conditions" (Figure 11a) or under "denaturing conditions" (Figure 11b).

Figures 12a-c show chemical, photochemical and enzymatic probing of dsDNA-Acr-T10-LysNH₂ complex. Complexes between Acr-T10-LysNH₂ and a ³²P-endlabeled DNA fragment containing a dA₁₀/dT₁₀ target sequence were probed by affinity photocleavage (Figure 12a, lanes 1-3; Figure 12b, lanes 1-3), photofootprinting (Figure 12a, lanes 5-6), potassium permanganate probing (Figure 12b, lanes 4-6) or probing by staphylococcus nuclease (Figure 12b, lanes 8-10) or by nuclease S₁ (Figure 12c). Either the A-strand (Figure 12a) or the T-strand (Figures 12b,c) was probed.

Figure 13 provides a procedure for the synthesis of 20 protected PNA synthons.

Figure 14 provides a procedure for the synthesis of a protected adenine monomer synthon.

Figure 15 provides a procedure for the synthesis of a protected guanine monomer synthon.

25 Figure 16 provides examples of PNA backbone alterations.

Figure 17 provides a procedure for synthesis of thymine monomer synthons with side chains corresponding to the normal amino acids.

Figures 18a and 18b provide procedures for synthesis of an aminopropyl analogue and a propionyl analogue, respectively, of a thymine monomer synthon.

Figure 19 provides a procedure for synthesis of an aminoethyl- β -alanine analogue of thymine monomer synthon.

Figure 20 shows a PAGE autoradiograph demonstrating that PNAs- T_{10} , - T_9 C and - T_8 C₂ bind to double stranded DNA with high sequence specificity.

Figure 21 shows a graph bas d n d nsit m tri scanning of PAGE autoradiographs d monstrating th kinetics of the binding of PNA- T_{10} to a double strand d target.

Figure 22 shows a graph based on densitometric scanning of PAGE autoradiographs demonstrating the thermal stabilities of PNAs of varying lengths bound to an A_{10}/T_{10} double strand d DNA target.

Figure 23 shows an electrophoretic gel staining demonstrating that restriction enzyme activity towards DNA is 10 inhibited when PNA is bound proximal to the restriction enzyme recognition site.

Figure 24 shows a PAGE autoradiograph demonstrating that $^{125}I\text{--labeled}$ PNA-T $_{10}$ binds to a complementary dA_{10} oligonucleotide.

15 Figure 25 shows a peptide nucleic acid according to the invention.

Figure 26 shows the direction of synthesis for a peptide nucleic acid according to the invention.

Figure 27 provides a test for the tosyl group as a 20 nitrogen protecting group in the synthesis of peptide nucl ic acids.

DETAILED DESCRIPTION OF THE INVENTION

In the oligonucleotide analogs and monomer synthons according to the invention, ligand L is primarily a naturally occurring nucleobase attached at the position found in nature, i.e., position 9 for adenine or guanine, and position 1 f r thymine or cytosine. Alternatively, L may be a non-naturally occurring nucleobase (nucleobase analog), another base-binding moiety, an aromatic moiety, (C₁-C₄)alkanoyl, hydroxy or even hydrogen. Some typical nucleobase ligands and illustrative synthetic ligands are shown in Figure 2. Furthermore, L can be a DNA intercalator, a reporter ligand such as, for example, a fluorophor, radio label, spin label, hapten, or a prot in-

In monomer synth ns, L may b blocked with protecting groups. This is illustrat d in Figure 4, wh r Pg is an

acid, a base or a hydr gen lytically or photochemically cleavable prot cting group such as, fr example, t-butoxycarb nyl (Boc), fluorenylmethyloxycarbonyl (Fmoc) or 2-nitrobenzyl (2Nb).

Linker A can be a wide variety of groups such as

-CR¹R²CO-, -CR¹R²CS-, -CR¹R²CSe-, -CR¹R²CNHR³-, -CR¹R²C=CH₂- and

-CR¹R²C=C(CH₃)₂-, where R¹, R² and R³ are as defined above.

Preferably, A is methylenecarbonyl (-CH₂CO-). Also, A can be
a longer chain moiety such as propanoyl, butancyl or

10 pentancyl, or corresponding derivative, wherein O is replaced
by another value of X or the chain is substituted with R¹R² or
is heterogenous, containing Y. Further, A can be a (C₂
C₆)alkylene chain, a (C₂-C₆)alkylene chain substituted with R¹R²
or can be heterogenous, containing Y. In certain cases, A can

15 just be a single bond.

In the preferred form of the invention, B is a nitrogen atom, thereby presenting the possibility of an achiral backbone. B can also be R^3N^4 , where R^3 is as defined above.

In the preferred form of the invention, C is -CR⁶R⁷-, 20 but can also be a two carbon unit, i.e. -CHR⁶CHR⁷- or -CR⁶R⁷CH₂-, where R⁶ and R⁷ are as defined above. R⁶ and R⁷ also can be a heteroaryl group such as, for example, pyrrolyl, furyl, thienyl, imidazolyl, pyridyl, pyrimidinyl, indolyl, or can be taken together to complete an alicyclic system such as, 25 for example, 1,2-cyclobutanediyl, 1,2-cyclopentanediyl or 1,2-cyclohexanediyl.

In the preferred form of the invention, E in the monomer synthon is COOH or an activated derivative thereof, and G in the oligomer is -CONR³-. As defined above, E may 30 also be CSOH, SOOH, SO₂OH or an activated derivative thereof, whereby G in the oligomer becomes -CSNR³-, -SONR³-and -SO₂NR³-, respectively. The activation may, for example, be achieved using an acid anhydride or an active ester derivative, wherein hydrogen in the groups represented by E is replaced by a 35 leaving group suited for generating the growing backbone.

The amin acids which f rm the backbone may be identical or different. We have found that those based on 2-

aminoethylglycine are especially well suited to the purpose of the invention.

In some cases it may be of interest to attach ligands at either terminus (Q, I) to modulate the binding characte5 ristics of the PNAs. Representative ligands include DNA intercalators which will improve dsDNA binding or basic groups, such as lysine or polylysine, which will strengthen the binding of PNA due to electrostatic interaction. To decrease negatively charged groups such as carboxy and sulf groups could be used. The design of the synthons furth r allows such other moieties to be located on non-terminal positions.

In a further aspect of the invention, the PNA oligomers are conjugated to low molecular effector ligands such as ligands having nuclease activity or alkylating activity r reporter ligands (fluorescent, spin labels, radioactive, protein recognition ligands, for example, biotin or haptens). In a further aspect of the invention, the PNAs are conjugated to peptides or proteins, where the peptides have signaling activity and the proteins are, for example, enzymes, transcription factors or antibodies. Also, the PNAs can be attached to water-soluble or water-insoluble polymers. In another aspect of the invention, the PNAs are conjugated to oligonucleotides or carbohydrates. When warranted, a PNA oligomer can be synthesized onto some moiety (e.g., a peptide chain, reporter, intercalator or other type of ligand-containing group) attached to a solid support.

Such conjugates can be used for gene modulation (e.g., gene targeted drugs), for diagnostics, for biotechnology, and 30 for scientific purposes.

As a further aspect of the invention, PNAs can be used to target RNA and ssDNA to produce both antisense-type gen regulating moieties and hybridization probes for the identification and purification of nucleic acids.

35 Furthermore, the PNAs can be modified in such a way that they can f rm triple helices with dsDNA. R agents that bind sequence-specifically to dsDNA hav applications as gene

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targeted drugs. These are foreseen as extremely useful drugs for treating diseases like cancer, AIDS and other virus infections, and may also prove effective for treatment of s me genetic diseases. Furthermore, these reagents may be used for 5 research and in diagnostics for detection and isolation of specific nucleic acids.

The triple helix principle is believed to be the only known principle in the art for sequence-specific recognition of dsDNA. However, triple helix formation is largely limited to recognition of homopurine-homopyrimidine sequences. Strand displacement is superior to triple helix recognition in that it allows for recognition of any sequence by use of the four natural bases. Also, in strand displacement recognition readily occurs at physiological conditions, that is, neutral ph, ambient (20-40 C) temperature and medium (100-150 mM) ionic strength.

Gene targeted drugs are designed with a nucleobase sequence (containing 10-20 units) complementary to the regulatory region (the promoter) of the target gene.

20 Therefore, upon administration of the drug, it binds to the promoter and block access thereto by RNA polymerase. Consequently, no mRNA, and thus no gene product (protein), is produced. If the target is within a vital gene for a virus, no viable virus particles will be produced. Alternatively, the target could be downstream from the promoter, causing the RNA polymerase to terminate at this position, thus forming a truncated mRNA/protein which is nonfunctional.

Sequence-specific recognition of ssDNA by base complementary hybridization can likewise be exploited to target specific genes and viruses. In this case, the target sequence is contained in the mRNA such that binding of the drug to the target hinders the action of ribosomes and, consequently, translation of the mRNA into protein. The peptide nucleic acids of the invention are superior to prior reagents in that they have significantly higher affinity for complementary ssDNA. Also, they p ssess n charge and water soluble, which should facilitate cellular uptake, and they contain amides of

non-biological amino acids, which should make th m biostable and r sistant to enzymatic degradati n by, for example, proteases.

Certain biochemical/biological properties of PNA 5 oligomers are illustrated by the following experiments.

1. Sequence discrimination at the dsDNA level (Example 63, Figure 20).

Using the S_1 -nuclease probing technique, the discrimination of binding of the T_{10} , $T_5CT_4(T_9C)$ & $T_2CT_2CT_4(T_8C_2)$ PNA 10 to the recognition sequences A_{10} , A_5GA_4 (A_9G) & $A_2GA_2GA_4$ (A_8G_2) cloned into the BamHI, SalI or PstI site of the plasmid pUC19 was analyzed. The results (Figure 20) show that the thre PNAs bind to their respective recognition sequences with the following relative efficiencies: PNA - T10: $A_{10} > A_9G >> A_8G_2$, 15 PNA - T_9C : $A_9G >> A_{10} - A_8G_2$, PNA - T_8C_2 : $A_8G_2 \ge A_9G >> A_{10}$. Thus at 37°C one mismatch out of ten gives reduced efficiency (5-10 times estimated) whereas two mismatches are not accepted.

- 2. Rinetics of PNA- T_{10} dsDNA strand displacement compl x formation (Example 66, Figure 21).
- Complex formation was probed by S_1 -nuclease at various times following mixing of PNA and 32 P-endlabeled dsDNA fragment (Figure 21).
- 3. Stability of PNA-dsDNA complex (Example 67, Figure 22)
 Complexes between PNA-T_n and ³²P-dsDNA (A₁₀/T₁₀) target
 25 were formed (60 min, 37°C). The complexes were then incubat d
 at the desired temperature in the presence of excess oligodA₁₀ for 10 min, cooled to RT and probed with KMnO₄. Th
 results (Figure 22) show that the thermal stability of the
 PNA-dsDNA complexes mirror that of the PNA oligonucleotide
 30 complexes in terms of "Tm".
 - 4. Inhibition of restriction enzyme cleavage by PNA (Example 65, Figure 23)

The plasmid construct, pT10, contains a dA_{10}/dT_{10} tract cloned into the BamHI site in pUC19. Thus, cleavage of pT10 35 with BamHI and PvuII results in two small DNA fragments of 211 and 111 bp, r sp ctively. In the presence of PNA- T_{10} , a 336 bp fragment is btained corr sponding t cleavage nly by

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PvuII (Figure 23). Thus cleavage by BamHI is inhibited by PNA bound proximal to the restriction enzyme site. The results also show that the PNA-dsDNA complex can be formed in 100% yield. Similar results were obtained using the pT8C2 plasmid and PNA-T8C2.

5. Binding of ¹²⁵I-labeled PNA to oligonucleotides (Example 63, Figure 24)

A Tyr-PNA-T₁₀-Lys-NH₂ was labeled with ¹²⁵I using Na¹²⁵I and chloramine-T and purified by HPLC. The ¹²⁵I-PNA-T₁₀ was 10 shown to bind to oligo-dA₁₀ by PAGE and autoradiography (Figure 24). The binding could be competed by excess denatured calf thymus DNA.

The sequence-specific recognition of dsDNA is illustrated by the binding of a PNA, consisting of 10 thymine substituted 2-aminoethylglycyl units, which C-terminates in a lysine amide and N-terminates in a complex 9-aminoacridine ligand (9-Acr¹-(Taeg)₁₀-Lys-NH₂, Figure 11a, 11b) to a dA₁₀/dT₁₀ target sequence. The target is contained in a 248 bp ³²P-end-labelled DNA-fragment.

- 20 Strand displacement was ascertained by the following type of experiments:
- 1) The 9-Acr¹ ligand (Figure 5), which is equipped with a 4-nitrobenzamido group to ensures cleavage of DNA upon irradiation, is expected only to cleave DNA in close proximity 25 to its binding site. Upon irradiation of the PNA with the above 248 bp DNA fragment, selective cleavage at the dA_{10}/dT_{10} sequence is observed (Figure 3a).
- In a so-called photofootprinting assay, where a synthetic diazo-linked acridine under irradiation cleaves DNA
 (except where the DNA is protected by said binding substance) upon interaction with DNA in the presence of a DNA-binding substance.

Such an experiment was performed with the above 248 bp dsDNA fragment, which showed clear protection against phosphocologies of the PNA binding site (Figure 3b).

3) In a similar type of experiment, the DNA-cleaving enzyme micrococcus nuclease, which is also hindered in its

action by most DNA-binding r agents, showed increased cleavage at the T_{10} -target (Figur 3c).

- 4) In yet another type of xperiment, the well-known high susceptibility of single strand thymine ligands (as 5 opposed to double strand thymine ligands) towards potassium permanganate oxidation was employed. Oxidation of the, 248 bp in the presence of the reagent showed only oxidation of the T₁₀-strand of the target (Figure 3b).
- 5) In a similar type of demonstration, the single 10 strand specificity of S_1 nuclease clearly showed that only th T_{10} -strand of the target was attacked (Figure 3d).

The very efficient binding of $(Taeg)_{10}$, $(Taeg)_{10}$ -Lys-NH₂ and Acr¹- $(Taeg)_{10}$ -Lys-NH₂ (Figures 11a. 11b) to the corresponding dA₁₀ was furthermore illustrated in two ways:

- 1. Ligand-oligonucleotide complexes will migrate slower than the naked oligonucleotide upon electrophoresis in polyacrylamide gels. Consequently, such experiments wer performed with Acr¹-(Taeg)₁₀-Lys-NH₂ and ³²P-end-labelled dA₁₀. This showed retarded migration under conditions where a normal dA₁₀/dT₁₀ duplex is stable, as well as under conditions where such a duplex is unstable (denaturing gel). A control experiment was performed with a mixture of Acr¹-(Taeg)₁₀-Lys-NH₂ and ³²P-end-labelled dT₁₀ which showed no retardation under the above conditions.
- 25 2. Upon formation of DNA duplexes (dsDNA) from single strand DNA, the extinction coefficient decreases (hypochromicity). Thus, the denaturing of DNA can be followed by measuring changes in the absorbance, for example, a a function of T_n, the temperature where 50% of a duplex has 30 disappeared to give single strands.

Duplexes were formed from the single-stranded oligodeoxyribonucleotides and the PNAs listed belw. Typically 0.3 OD₂₆₀ of the T-rich strand was hybridized with 1 equivalent of the other strand by heating to 90 C for 5 min, 35 cooling to room temperature and kept for 30 min and finally st red in a refrigerator at 5 C for at least 30 min. The buffers us d were all 10 mM in phosphat and 1 mM in EDTA.

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The low salt buffer contained no sodium chloride, whereas the medium salt buffer c ntained 140 mM NaCl and the high salt buffer 500 mM NaCl. The pH of all the buffers was 7.2. The melting temperature of the hybrids were determined on a 5 Gilford Response apparatus. The following extinction coefficients were used A: 15.4 ml/µmol'cm; T: 8.8; G: 11.7 and C: 7.3 for both normal oligonucleotides and PNA. The melting curves were recorded in steps of 0.5 C/min. The T_a were determined from the maximum of the 1st derivative of the plot 10 of A₂₆₀ vs temperature.

List of oligodeoxyribonucleotides:

- 1. 5'-AAA-AAA-AA
- 2. 5'-AAA-AAA-AAA-A
- 15 3. 5'-TTT-TTT-T
 - 4. 5'-AAA-AAG-AAA-A
 - 5. 5'-AAG-AAG-AAA-A
 - 6. 5'-AAA-AGA-AAA-A
 - 7. 5'-AAA-AGA-AGA-A
- 20 8. 5'-TTT-TCT-TTT-T
 - 9. 5'-TTT-TCT-TCT-T
 - 10. 5'-TTT-TTC-TTT-T
 - 11. 5'-TTT-TTC-TTC-T
 - 12. 5'-TTC-TTC-TTT-T
- 25 13. 5'-TTT-TTT-TTT-TTT
 - 14. 5'-AAA-AAA-AAA-AAA

List of PNAs

- a. TTT-TTT-TTT-Lys-NH,
- 30 b. TTT-TTT-TT-Lys-NH,
 - c. TTT-TTC-TTT-T-Lys-NH2
 - d. TTC-TTC-TTT-T-Lys-NH,
 - e. ACT-TTT-TTT-T-Lys-NH,
 - f. AC-TTT-TTT-TT-Lys-NH₂

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	Oligo/PNA	Low Salt	Medium Salt	High Salt
	1 + b	56.0	51.5	50.0
	_ 2+a	73.0	72.5	73.0
	2+c		41.5 and 52.0°	
5	2+e	84.5	86.0	-90
	2+1		74	
	4+8	60.0	59.0	61.5
L	4+c -	74.5	72.0	72.5
	4+f		62.0	
10	5+a		47.0	
	5+c		57.5	
	5+f		46.5	
L	7+8		46.0	-
	7+c		58.0	
15	7+f		43.5	
	7+12		23.0	
	13+14		39.0	

* = Two distinct melting temperatures are seen, indicating local melting before complete denaturation.

The hybrid formed between RNA-A (poly rA) and PNA- T_{10} -Lys-NH₂ melts at such high temperature that it cannot b measured (>90 C). But specific hybridization is demonstrated by the large drop in λ_{260} by mixing with RNA-A but not G,C and U. The experiment is done by mixing 1 ml of a solution of th PNA and 1 ml of a solution the RNA, each with $\lambda_{260} = 0.6$, and then measure the absorbance at 260 nm. Thereafter the sample is heated to 90 C for 5 min, cooled to room temperature and 30 kept at this temperature for 30 minutes and finally stored at 5C for 30 min.

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RNA	PNA	A ₂₆₀ Bef re Mixing	A ₂₆₀ After Mixing	A ₂₆₀ After Mixing and Heating
RNA-A	PNA-Tulys-NH,	0.600	0.389	0.360
RNA-U	PNA-T ₁₀ -lys-NH,	0.600	0.538	0.528
RNA-G	PNA-T ₁₀ -lys-NH,	0.600	0.514	0.517
RNA-C	PNA-T ₁₀ -lys-NH,	0.600	0.540	0.532

From the above measurements the following conclusions can be made. There is base stacking, since a melting curve is observed. The PNA-DNA hybrid is more stable than a normal 10 DNA-DNA hybrid, and the PNA-RNA is even more stable. Mismatches cause significant drops in the Tm-value, whether the mispaired base is in the DNA or in the PNA-strand. The Tm-value is only slightly dependent on ionic strength, as opposed to normal oligonucleotides.

The synthesis of the PNAs according to the invention is discussed in detail in the following, where Figure 1 illustrates one of the preferred PNA examples and compares its structure to that of a complementary DNA.

The principle of anchoring molecules onto a solid

20 Synthesis of PNA Oligomers and Polymers

present invention (Figure 8).

during chemical transformations, is known as Solid-Phase Synthesis or Merrifield Synthesis (see, e.g., Merrifield, J. 25 Am. Chem. Soc., 1963, 85, 2149 and Science, 1986, 232, 341). Established methods for the stepwise or fragmentwise solid-phase assembly of amino acids into peptides normally employ a beaded matrix of slightly cross-linked styrene-divinylbenzene copolymer, the cross-linked copolymer having been formed by the pearl polymerization of styrene monomer to which has been added a mixture of divinylbenzenes. A level of 1-2% cross-linking is usually employed. Such a matrix also can be us d in solid-phase PNA synthesis in accordance with the

matrix, which helps in accounting for intermediate products

Concerning the initial functionalization of the s lid phase, more than fifty methods hav been d scribed in connection with traditional solid-phas p ptide synthesis (see, e.g., Barany and Merrifield in "The Peptides" Vol. 2, 5 Academic Press, New York, 1979, pp. 1-284, and Stewart and Young, "Solid Phase Peptide Synthesis", 2nd Ed., Pierc Chemical Company, Illinois, Reactions for the 1984). introduction of chloromethyl functionality (Merrifield resin; via a chloromethyl methyl ether/SnCl, reaction), aminomethyl 10 functionality (via an N-hydroxymethylphthalimide reacti n; see, Mitchell, et al., Tetrahedron Lett., 1976, 3795), and benzhydrylamino functionality (Pietta, et al., J. Chem. Soc., 1970, 650) are the most widely applied. Regardless of its nature, the purpose of the functionality is normally to f rm 15 an anchoring linkage between the copolymer solid support and the C-terminus of the first amino acid to be coupled to the solid support. As will be recognized, anchoring linkages also . can be formed between the solid support and the amino acid N-It is generally convenient to express the terminus. 20 "concentration" of a functional group in terms of millimoles per gram (mmol/g). Other reactive functionalities which hav been initially introduced include 4-methylbenzhydrylamino and 4-methoxybenzhydrylamino. All of these established methods are in principle useful within the context of the present in-25 vention. Preferred methods for PNA synthesis employ aminomethyl as the initial functionality, in that aminomethyl is particularly advantageous with respect to the incorporati n of "spacer" or "handle" groups, owing to the reactivity of the amino group of the aminomethyl functionality with respect to 30 the essentially quantitative formation of amide bonds to a carboxylic acid group at one end of the spacer-forming reagent. A vast number of relevant spacer- or handle-forming bifunctional reagents have been described (see, Barany, t al., Int. J. Peptide Protein Res., 1987, 30, 705), especially 35 reagents which are reactive towards amino groups such as found in the aminomethyl function. Representative bifunctional r ag nts include 4-(haloalkyl)aryl-lower alkan ic acids such

4-(bromomethyl)ph nylacetic acid, Boc-aminoacy1-4-(oxymethyl)aryl-lower alkanoic acids such as Boc-aminoacyl-4-(oxymethyl) phenylacetic acid, N-Boc-p-acylbenzhydrylamines such as N-Boc-p-glutaroylbenzhydrylamine, N-Boc-4'-lower 5 alkyl-p-acylbenzhydrylamines such as N-Boc-4'-methyl-pglutaroylbenzhydrylamine, N-Boc-4'-lower alkoxy-p-acylbenzhydrylamines such as N-Boc-4'-methoxy-p-glutaroyl-benzhydrylamine, and 4-hydroxymethylphenoxyacetic acid. One type of spacer group particularly relevant within the context f 10 the present invention is the phenylacetamidomethyl (Pam) handle (Mitchell and Merrifield, J. Org. Chem., 1976, 41, 2015) which, deriving from the electron withdrawing effect of the 4-phenylacetamidomethyl group, is about 100 times more stable than the classical benzyl ester linkage towards the 15 Boc-amino deprotection reagent trifluoroacetic acid (TFA).

Certain functionalities (e.g., benzhydrylamino, 4-methylbenzhydrylamino and 4-methoxybenzhydrylamino) which may be incorporated for the purpose of cleavage of a synthesized PNA chain from the solid support such that the C-terminal of the PNA chain is in amide form, require no introduction of a spacer group. Any such functionality may advantageously be employed in the context of the present invention.

An alternative strategy concerning the introduction of spacer or handle groups is the so-called "preformed handl"

25 strategy (see, Tam, et al., Synthesis, 1979, 955-957), which offers complete control over coupling of the first amino acid, and excludes the possibility of complications arising from the presence of undesired functional groups not related to the peptide or PNA synthesis. In this strategy, spacer or handle groups, of the same type as described above, are reacted with the first amino acid desired to be bound to the solid supp rt, the amino acid being N-protected and optionally protected at the other side-chains which are not relevant with respect to the growth of the desired PNA chain. Thus, in those cases in which a spacer or handle group is desirable, the first amino acid to be coupled to the free reactive end of a spacer group which has been

bound to the initially introduced functi nality (for example, an aminomethyl group) or can b react d with the spacer-forming reagent. The space-f rming reagent is then react d with the initially introduced functionality. Other useful anchoring schemes include the "multidetachable" resins (Tam, et al., Tetrahedron Lett., 1979, 4935 and J. Am. Chem, Soc., 1980, 102, 611; Tam, J. Org. Chem., 1985, 50, 5291), which provide more than one mode of release and thereby allow more flexibility in synthetic design.

Suitable choices for N-protection are the tert-10 butyloxycarbonyl (Boc) group (Carpino, J. Am. Chem. Soc., 1957, 79, 4427; McKay, et al., J. Am. Chem. Soc., 1957, 79, 4686; Anderson, et al., J. Am. Chem. Soc., 1957, 79, 6180) normally in combination with benzyl-based groups for the 15 protection of side chains, and the 9-fluorenylmethyloxycarbonyl (Fmoc) group (Carpino, et al., J. Am. Chem. Soc., 1970, 92, 5748 and J. Org. Chem., 1972, 37, 3404), normally in combination with tert-butyl (tBu) for the protection of any side chains, although a number of other possibilities exist 20 which are well known in conventional solid-phase peptid Thus, a wide range of other useful amin synthesis. protecting groups exist, some of which are Adoc (Hass, et al., J. Am. Chem. Soc., 1966, 88, 1988), Bpoc (Sieber, Helv. Ch m. Acta., 1968, 51, 614), Mcb (Brady, et al., J. Org. Chem., 25 1977, 42, 143), Bic (Kemp, et al., Tetrahedron, 1975, 4624), the o-nitrophenylsulfenyl (Nps) (Zervas, et al., J. Am. Ch m. Soc., 1963, 85, 3660), and the dithiasuccinoyl (Dts) (Barany, et al., J. Am. Chem. Soc., 1977, 99, 7363). protecting groups, particularly those based on the widely-us d 30 urethane functionality, successfully prohibit racemization (mediated by tautomerization of the readily formed oxazolinone (azlactone) intermediates (Goodman, et al., J. Am. Chem. Soc., 1964, 86, 2918)) during the coupling of most α-amino acids. In addition to such amino protecting groups, a whole range of 35 oth rwis "w rthl ss" nonur thane-type of amino protecting groups are applicable when assembling PNA m lecules,

especially th se built fr m achiral units. Thus, not only the above-mentioned amino protecting groups (or those derived from any of these groups) are useful within the context of the present invention, but virtually any amino protecting group 5 which largely fulfills the following requirements: (1) stability to mild acids (not significantly attacked by carboxyl groups); (2) stability to mild bases or nucleophiles (not significantly attacked by the amino group in question); (3) resistance to acylation (not significantly attacked by 10 activated amino acids). Additionally: (4) the protecting group must be close to quantitatively removable, without serious side reactions, and (5) the optical integrity, if any, of the incoming amino acid should preferably be highly preserved upon coupling. Finally, the choice of side-chain 15 protecting groups, in general, depends on the choice of the amino protecting group, since the protection of side-chain functionalities must withstand the conditions of the repeated amino deprotection cycles. This is true whether the overall strategy for chemically assembling PNA molecules relies on, 20 for example, differential acid stability of amino and sidechain protecting groups (such as is the case for the abovementioned "Boc-benzyl" approach) or employs an orthogonal, that is, chemoselective, protection scheme (such as is the case for the above-mentioned "Fmoc-tBu" approach),

25 Following coupling of the first amino acid, the next stage of solid-phase synthesis is the systematic elaboration of the desired PNA chain. This elaboration involves repeated deprotection/coupling cycles. The temporary protecting group, such as a Boc or Fmoc group, on the last-coupled amino acid 30 is quantitatively removed by a suitable treatment, for example, by acidolysis, such as with trifluoroacetic acid, in the case of Boc, or by base treatment, such as with piperidine, in the case of Fmoc, so as to liberate the N-terminal amine function.

35 The next desired N-protected amino acid is then coupled to the N-terminal f the last-c upled amino acid. This coupling of the C-terminal of an amino acid with the N-

terminal f the last-coupl d amin acid can be achieved in several ways. For xample, the carboxyl group f the inc ming amino acid can be reacted directly with the N-t rminal f the last-coupled amino acid with the assistance of a condensation 5 reagent such as, for example, dicyclohexylcarbodiimide (DCC) (Sheehan & Hass, et al., J. Am. Chem. Soc., 1955, 77, 1067) and diisoproplycarbodiimide (DIC) (Sraantakis et al., Biochem. biophys. res. Commun., 1976, 73, 336) or derivatives thereof. Alternatively, it can be bound by providing the incoming amino 10 acid in a form with the carboxyl group activated by any of several methods, including the initial formation of an active ester derivative such as a 2,4,5-trichlorophenyl ester (Pless, et al., Helv. Chim. Acta, 1963, 46, 1609), a phthalimido ester (Nefkens, et al., J. Am. Chem. Soc., 1961, 83, 1263), a 15 pentachlorophenyl ester (Kupryszewski, Rocz. Chem., 1961, 35, 595), a pentafluorophenyl ester (Kovacs, et al., J. Am. Chem. Soc., 1963, 85, 183), an o-nitrophenyl ester (Bodanzsky, Nature, 1955, 175, 685), an imidazole ester (Li, et al., J. Am. Chem. Soc., 1970, 92, 7608), and a 3-hydroxy-4-oxo-3,4-20 dihydroquinazoline (Dhbt-OH) ester (Konig, et al., Chem. Ber., 1973, 103, 2024 and 2034), or the initial formation of an anhydride such as a symmetrical anhydride (Wieland, et al., Angew. Chem., Int. Ed. Engl., 1971, 10, 336). Benzotriazolyl N-oxytrisdimethylaminophosphonium hexafluorophosphate (BOP), 25 "Castro's reagent" (see, e.g., Rivaille, et al., Tetrahedron, 1980, 36, 3413) is recommended when assembling PNA molecules containing secondary amino groups. Finally, activated PNA monomers analogous to the recently-reported amino acid fluorides (Carpino, J. Am. Chem. Soc., 1990, 112, 9651) h ld 30 considerable promise to be used in PNA synthesis as well.

Following assembly of the desired PNA chain, including protecting groups, the next step will normally be deprotect in of the amino acid moieties of the PNA chain and cleavage of the synthesized PNA from the solid support. These processes can take place substantially simultaneously, thereby providing the free PNA m 1 cule in the d sir d form. Alternatively, in cases in which condensation of two separately synth sized PNA cases in which condensation of two separately synth sized PNA

chains is to be carried ut, it is possible by choosing a suitable spacer group at the start of the synthesis t cleave the desired PNA chains from their respective solid supports (both peptide chains still incorporating their side-chain protecting groups) and finally removing the side-chain protecting groups after, for example, coupling the two side-chain protected peptide chains to form a longer PNA chain.

In the above-mentioned "Boc-benzyl" protection scheme, the final deprotection of side-chains and release of the PNA 10 molecule from the solid support is most often carried out by the use of strong acids such as anhydrous HF (Sakakibara, t al., Bull. Chem. Soc. Jpn., 1965, 38, 4921), boron tris (trifluoroacetate) (Pless, et al., Helv. Chim. Acta, 1973, 46, 1609), and sulfonic acids such as trifluoromethanesulfonic 15 acid and methanesulfonic acid (Yajima, et al., J. Chem. Soc., Chem. Comm., 1974, 107). This conventional strong acid (e.g., anhydrous HF) deprotection method, produces very reactive carbocations that may lead to alkylation and acylation of sensitive residues in the PNA chain. Such side-reactions are 20 only partly avoided by the presence of scavengers such as anisole, phenol, dimethyl sulfide, and mercaptoethanol and, therefore, the sulfide-assisted acidolytic Su2 deprotection method (Tam, et al., J. Am. Chem. Soc., 1983, 105, 6442 and J. Am. Chem. Soc., 1986, 108, 5242), the so-called "low", 25 which removes the precursors of harmful carbocations to form inert sulfonium salts, is frequently employed in peptide and PNA synthesis, either solely or in combination with "high" methods. Less frequently, in special cases, other methods used for deprotection and/or final cleavage of the PNA-solid 30 support bond are, for example, such methods as base-catalyzed -alcoholysis (Barton, et al., J. Am. Chem. Soc., 1973, 95, 4501), and ammonolysis as well as hydrazinolysis (Bodanszky, et al., Chem. Ind., 1964 1423), hydrogenolysis (Jones, Tetrahedron Lett. 1977 2853 and Schlatter, et al., Tetrahedron 35 L tt. 1977 2861)), and photolysis (Rich and Gurwara, J. Am. Chem. Soc., 1975 97, 1575)).

Finally, in contrast with the hemical synth sis of "normal" peptides, stepwise chain building f achiral PNAs such as those based n aminoethylglycyl backbon units can start either from the N-terminus or the C-terminus, because 5 the coupling reactions are free of racemization. Those skilled in the art will recognize that whereas syntheses commencing at the C-terminus typically employ protected amine groups and free or activated acid groups, syntheses commencing at the N-terminus typically employ protected acid groups and 10 free or activated amine groups.

Based on the recognition that most operations ar identical in the synthetic cycles of solid-phase peptide synthesis (as is also the case for solid-phase PNA synthesis), a new matrix, PEPS, was recently introduced (Berg, et al., J. 15 Am. Chem. Soc., 1989, 111, 8024 and International Patent Application WO 90/02749) to facilitate the preparation f large numbers of peptides. This matrix is comprised of a polyethylene (PE) film with pendant long-chain polystyrene (PS) grafts (molecular weight on the order of 10°). 20 loading capacity of the film is as high as that of a beaded matrix, but PEPS has the additional flexibility to suit multiple syntheses simultaneously. Thus. configuration for solid-phase peptide synthesis, the PEPS film is fashioned in the form of discrete, labeled sheets, each 25 serving as an individual compartment. During all the identical steps of the synthetic cycles, the sheets are kept together in a single reaction vessel to permit concurrent preparation of a multitude of peptides at a rate close to that of a single peptide by conventional methods. It was reasoned 30 that the PEPS film support, comprising linker or spacer groups adapted to the particular chemistry in question, should be - particularly valuable in the synthesis of multiple PNA molecules, these being conceptually simple to synthesize sinc only four different reaction compartments are normally 35 required, one for each of the four "pseudo-nucleotide" units. Thus, the PEPS film support has been successfully tested in a number f PNA synth ses carried out in a parallel and

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substantially simultaneous fashion. The yield and quality of the pr ducts brained from PEPS were comparable to those obtained by using the traditional polystyrene beaded support. Also, experiments with other geometries of the PEPS polymer such as, for example, non-woven felt, knitted net, sticks or microwellplates have not indicated any limitations of the synthetic efficacy.

Two other methods proposed for the simultaneous synthesis of large numbers of peptides also apply to the 10 preparation of multiple, different PNA molecules. The first of these methods (Geysen, et al., Proc. Natl. Acad. Sci. USA, 1984, 81, 3998) utilizes acrylic acid-grafted polyethylen rods and 96-microtiter wells to immobilize the growing peptide chains and to perform the compartmentalized synthesis. While 15 highly effective, the method is only applicable on a microgram scale. The second method (Houghten, Proc. Natl. Acad. Sci. USA, 1985, 82, 5131) utilizes a "tea bag" containing traditionally-used polymer beads. Other relevant proposals for multiple peptide or PNA synthesis in the context of the 20 present invention include the simultaneous use of two different supports with different densities (Tregear, in "Chemistry and Biology of Peptides", J. Meienhofer, ed., Ann Arbor Sci. Publ., Ann Arbor, 1972 pp. 175-178), combining of reaction vessels via a manifold (Gorman, Anal. Biochem., 1984, 25 136, 397), multicolumn solid-phase synthesis (e.g. Krchnak, et al., Int. J. Peptide Protein Res., 1989, 33, 209), and Holm and Meldal, in "Proceedings of the 20th European Peptide Symposium", G. Jung and E. Bayer, eds., Walter de Gruyter & Co., Berlin, 1989 pp. 208-210), and the use of cellulose paper 30 (Eichler, et al., Collect. Czech. Chem. Commun., 1989, 54, 1746).

While the conventional cross-linked styrene/divinylbenzene copolymer matrix and the PEPS support are presently preferred in the context of solid-phase PNA synthesis, a non-limiting list of examples of solid supports which may be of relevance are: (1) Particles based upon copolymers of dimethylacrylamide cross-linked with N,N'-

bisacryloylethyl nediamine, including a known amount of Nrtbutoxycarb nyl-beta-alanyl-N'acryloylhexamethylen diamin . Sev ral spacer mol cules are typically added via the beta alanyl group, followed thereafter 5 by the amino acid residue subunits. Also, the beta alanylcontaining monomer can be replaced with an acryloyl sarcosine monomer during polymerization to form resin beads. polymerization is followed by reaction of the beads with ethylenediamine to form resin particles that contain primary 10 amines as the covalently linked functionality. polyacrylamide-based supports are relatively more hydrophilic than are the polystyrene-based supports and are usually us d with polar aprotic solvents including dimethylformamide, dimethylacetamide, N-methylpyrrolidone and the like (s 15 Atherton, et al., J. Am. Chem. Soc., 1975, 97, 6584, Bioorg. Chem. 1979, 8, 351), and J.C.S. Perkin I 538 (1981)); (2) a second group of solid supports is based on silica-containing particles such as porous glass beads and silica gel. example is the reaction product of trichloro-[3-(4-20 chloromethyl)phenyl]propylsilane and porous glass beads (see Parr and Grohmann, Angew. Chem. Internal. Ed. 1972, 11, 314) sold under the trademark "PORASIL E" by Waters Associates, Framingham, MA, USA. Similarly, a mono ester of 1,4-dihydroxymethylbenzene and silica (sold under the trademark "BIOPAK" 25 by Waters Associates) has been reported to be useful (se Bayer and Jung, Tetrahedron Lett., 1970, 4503); (3) a third general type of useful solid supports can be termed composites in that they contain two major ingredients: a resin and another material that is also substantially inert to th 30 organic synthesis reaction conditions employed. One exemplary composite (see Scott, et al., J. Chrom. Sci., 1971, 9, 577) utilized glass particles coated with a hydrophobic, crosslinked styrene polymer containing reactive chloromethyl groups, and was supplied by Northgate Laboratories, Inc., of 35 Hamden, CT, USA. Another exemplary composite contains a c re f flu rinated ethylene p lymer onto which has been grafted p lystyren (see Kent and Merrifi ld, Isra l J. Chem. 1978,

17, 243) and van Rietschoten in "P ptides 1974", Y. Wolman, Ed., Wiley and Sons, New York, 1975, pp. 113-116); and (4) contiguous solid supports other than PEPS, such as cotton sheets (Lebl and Eichler, Peptide Res. 1989, 2, 232) and hydroxypropylacrylate-coated polypropylene membranes (Daniels, et al., Tetrahedron Lett. 1989, 4345), are suited for PNA synthesis as well.

Whether manually or automatically operated, solid-phase PNA synthesis in the context of the present invention is normally performed batchwise. However, most of the syntheses may equally well be carried out in the continuous-flow mode, where the support is packed into columns (Bayer, et al., Tetrahedron Lett., 1970, 4503 and Scott, et al., J. Chromatogr. Sci., 1971, 9, 577). With respect to continuous-flow solid-phase synthesis, the rigid poly(dimethylacrylamide)-Kieselguhr support (Atherton, et al., J. Chem. Soc. Ch m. Commun., 1981, 1151) appears to be particularly successful, but another valuable configuration concerns the one worked out for the standard copoly(styrene-1%-divinylbenzene) support (Krchnak, et al., Tetrahedron Lett., 1987, 4469).

While the solid-phase technique is presently preferred in the context of PNA synthesis, other methodologies or combinations thereof, for example, in combination with the solid-phase technique, apply as well: (1) the classical 25 solution-phase methods for peptide synthesis (e.g., Bodanszky, "Principles of Peptide Synthesis", Springer-Verlag, Berlin-New York 1984), either by stepwise assembly or by segment/fragment condensation, are of particular relevance when considering especially large scale productions (gram, kilogram, and even 30 tons) of PNA compounds; (2) the so-called "liquid-phase" strategy, which utilizes soluble polymeric supports such as linear polystyrene (Shemyakin, et al., Tetrahedron Lett., 1965, 2323) and polyethylene glycol (PEG) (Mutter and Bayer, Angew. Chem., Int. Ed. Engl., 1974, 13, 88), is useful; (3) 35 random polymerization (s e, .g., Odian, "Principles of Polymerization", McGraw-Hill, New York (1970)) yielding

mixtures of many m lecular weights ("polydisp rse") peptide or PNA molecules are particularly relevant for purposes such as screening for antiviral effects; (4) a t chnique based on the use of polymer-supported amino acid active esters 5 (Fridkin, et al., J. Am. Chem. Soc., 1965, 87, 4646), sometimes referred to as "inverse Merrifield synthesis" or "polymeric reagent synthesis", offers the advantage of isolation and purification of intermediate products, and may thus provide a particularly suitable method for the synthesis 10 of medium-sized, optionally protected, PNA molecules, that can subsequently be used for fragment condensation into larger PNA molecules; (5) it is envisaged that PNA molecules may be assembled enzymatically by enzymes such as proteases or derivatives thereof with novel specificities (obtained, f r 15 example, by artificial means such as protein engineering). Also, one can envision the development of "PNA ligases" f r the condensation of a number of PNA fragments into very large PNA molecules; (6) since antibodies can be generated t virtually any molecule of interest, the recently develop d 20 catalytic antibodies (abzymes), discovered simultaneously by the groups of Lerner (Tramantano, et al., Science, 1986, 234, 1566) and of Schultz (Pollack, et al., Science, 1986, 234, 1570), should also be considered as potential candidates for assembling PNA molecules. Thus, there has been considerable in producing abzymes catalyzing acyl-transfer 25 success reactions (see for example Shokat, et al., Nature, 1989, 338, 269) and references therein). Finally, completely artificial enzymes, very recently pioneered by Stewart's group (Hahn, t al., Science, 1990, 248, 1544), may be developed to suit PNA The design of generally applicable enzymes, 30 synthesis. ligases, and catalytic antibodies, capable of mediating specific coupling reactions, should be more readily achieved for PNA synthesis than for "normal" peptide synthesis since PNA molecules will often be comprised of only four differ nt 35 amino acids (one for each of the four native nucleobases) as compared t the twenty natural by occurring (proteinogenic) amino acids constituting peptid s. In conclusion, no single strategy may be wholly suitable for the synthesis of a specific PNA molecule, and therefore, sometimes a combination of methods may work best.

The present invention also is directed to therapeutic 5 or prophylactic uses for peptide nucleic acids. therapeutic and prophylactic targets include herpes simpl x virus (HSV). human papillomavirus (HPV), immunodeficiency virus (HIV), candidia albicans, influenza virus, cytomegalovirus (CMV), intracellular adhesion molecul s 10 (ICAM), 5-lipoxygenase (5-LO), phospholipase A₂ (PLA₂), protein kinase C (PKC), and RAS oncogene. Potential applications of such targeting include treatments for ocular, labial, genital, and systemic herpes simplex I and II infections; genital warts; cervical cancer; common warts; 15 Kaposi's sarcoma; AIDS; skin and systemic fungal infections; flu; pneumonia; retinitis and pneumonitis in immunosuppressed patients; mononucleosis; ocular, skin and inflammation; cardiovascular disease; cancer; psoriasis; cardiovascular collapse; cardiac infarction; 20 gastrointestinal disease; kidney disease; arthritis; osteoarthritis; acute pancreatitis; septic shock; and Crohn's disease.

For therapeutic or prophylactic treatment, the peptide nucleic acids of the invention can be formulated in a 25 pharmaceutical composition, which may include carriers, thickeners, diluents, buffers, preservatives, surface active agents and the like. Pharmaceutical compositions may also include one or more active ingredients such as antimicrobial agents, antiinflammatory agents, anesthetics, and the like in 30 addition to peptide nucleic acid.

The pharmaceutical composition may be administered in a number of ways depending on whether local or systemic treatment is desired, and on the area to be treated. Administration may be done topically (including ophthalmically, vaginally, r ctally, intranasally), orally, by inhalation, or parenterally, for xample by intravenous

drip or subcutaneous, intraperit neal r intramuscular injection.

Formulations for topical administration may include ointments, lotions, creams, gels, drops, suppositories, 5 sprays, liquids and powders. Conventional pharmaceutical carriers, aqueous, powder or oily bases, thickeners and the like may be necessary or desirable. Coated condoms may also be useful.

Compositions for oral administration include powders or granules, suspensions or solutions in water or non-aqueous media, capsules, sachets, or tablets. Thickeners, flavorings, diluents, emulsifiers, dispersing aids or binders may be desirable.

Formulations for parenteral administration may includ
15 sterile aqueous solutions which may also contain buffers,
diluents and other suitable additives.

Dosing is dependent on severity and responsiveness of the condition to be treated, but will normally be one or more doses per day, with course of treatment lasting from several 20 days to several months or until a cure is effected or a diminution of disease state is achieved. Persons of ordinary skill can easily determine optimum dosages, dosing methodologies and repetition rates.

Treatments of this type can be practiced one a variety
of organisms ranging from unicellular prokaryotic and eukaryotic organisms to multicellular eukaryotic organisms. Any
organism that utilizes DNA-RNA transcription or RNA-protein
translation as a fundamental part of its hereditary, metabolic
or cellular control is susceptible to therapeutic and/or
prophylactic treatment in accordance with the invention.
Seemingly diverse organisms such as bacteria, yeast, protozoa,
algae, all plants and all higher animal forms, including warmblooded animals, can be treated. Further, since each cell of
multicellular eukaryotes can be treated since they include
both DNA-RNA transcription and RNA-protein translation as
integral parts of their cellular activity. Furtherm re, many
of the organelles (.g., mitochondria and chloroplasts) of

eukaryotic cells also include transcription and translation mechanisms. Thus, single cells, cellular populations or organelles can also be included within the definition of organisms that can be treated with therapeutic or diagnostic phosphorothicate oligonucleotides. As used herein, therapeutics is meant to include the eradication of a disease state, by killing an organism or by control of erratic r harmful cellular growth or expression.

The present invention also pertains to the advantage us 10 use of PNA molecules in solid-phase biochemistry (see, e.g., "Solid-Phase Biochemistry - Analytical and Synthetic Aspects", W. H. Scouten, ed., John Wiley & Sons, New York, 1983), notably solid-phase biosystems, especially bioassays or solidphase techniques which concerns diagnostic detection/quanti-15 tation or affinity purification of complementary nucleic acids (see, e.g., "Affinity Chromatography - A Practical Approach", P. D. G. Dean, W. S. Johnson and F. A. Middle, eds., IRL Press Ltd., Oxford 1986; "Nucleic Acid Hybridization - A Practical Approach", B. D. Harnes and S. J. Higgins, IRL Press Ltd., Present day methods for performing such 20 Oxford 1987). bioassays or purification techniques almost exclusively utilize "normal" or slightly modified oligonucleotides either physically adsorbed or bound through a substantially permanent covalent anchoring linkage to beaded solid supports such as 25 cellulose, glass beads, including those with controlled porosity (Mizutani, et al., J. Chromatogr., 1986, 356, 202), "Sephadex", "Sepharose", agarose, polyacrylamide, porous particulate alumina, hydroxyalkyl methacrylate gels, diolbonded silica, porous ceramics, or contiguous-materials such 30 as filter discs of nylon and nitrocellulose. One example employed the chemical synthesis of oligo-dT om cellulose beads for the affinity isolation of poly A tail containing mRNA (Gilham in "Methods in Enzymology," L. Grossmann and K. Moldave, eds., vol. 21, part D, page 191, Academic Press, New 35 York and London, 1971). All the above-menti ned methods are applicable within the context of the present inventi n. However, when possible, c valent linkage is preferred over the

physical adsorption of the molecules in question, since the disadvantag that s me of th latt r appr ach has th immobilized m lecules can be washed ut (desorbed) during the hybridization or affinity process. There is, thus, littl 5 control of the extent to which a species adsorbed on the surface of the support material is lost during the yarious treatments to which the support is subjected in the course of the bioassay/purification procedure. The severity of this problem will, of course, depend to a large extent on the rate 10 at which equilibrium between adsorbed and "free" species is established. In certain cases it may be virtually impossible to perform a quantitative assay with acceptable accuracy and/or reproducibility. Loss of adsorbed species during treatment of the support with body fluids, aqueous reagents 15 or washing media will, in general, be expected to be m st pronounced for species of relatively low molecular weight. In contrast with oligonucleotides, PNA molecules are easier to attach onto solid supports because they contain str ng nucleophilic and/or electrophilic centers. In addition, th 20 direct assembly of oligonucleotides onto solid supports suffers from an extremely low loading of the immobilized molecule, mainly due to the low surface capacity of the materials that allow the successful use of the state-of-theart phosphoramidite chemistry for the construction of oligo-25 nucleotides. (Beaucage and Caruthers, Tetrahedron Lett., 1981, 22, 1859; Caruthers, Science, 1985, 232, 281). suffers from the fact that by using the alternative phosphite triester method (Letsinger and Mahadevan, J. Am. Chem. Soc. 1976, 98, 3655), which is suited for solid supports with a 30 high surface/loading capacity, only relatively short oligonucleotides can be obtained. As for conventional solid-phase peptide synthesis, however, the latter supports are excellent materials for building up immobilized PNA molecules (the sidechain protecting groups are removed from the synthesized PNA 35 chain without cleaving the anchoring linkage holding the chain to the s lid support). Thus, PNA species benefit from th above-described solid-phase t chniques with resp ct to the

much higher (and still sequence-specific) binding affinity f r complementary nucleic acids and from the additional unique sequence-specific recognition of (and strong binding to) nucleic acids present in double-stranded structures. 5 also can be loaded onto solid supports in large amounts, thus further increasing the sensitivity/capacity of the solid-phase technique. Further, certain types of studies concerning the use of PNA in solid-phase biochemistry can be approached, facilitated, or greatly accelerated by use of the recently-10 reported "light-directed, spatially addressable, parallel chemical synthesis" technology (Fodor, et al., Science, 1991, 251, 767), a technique that combines solid-phase chemistry and photolithography to produce thousands of highly diverse, but identifiable, permanently immobilized compounds (such as 15 peptides) in a substantially simultaneous way.

Additional objects, advantages, and novel features f this invention will become apparent to those skilled in the art upon examination of the following examples thereof, which 20 are not intended to be limiting.

Synthesis of monomeric building blocks

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The monomers preferably are synthesized by the general scheme outlined in Figure 13. This involves preparation of 25 either the methyl or ethyl ester of (Bocaminoethyl)glycin , by a protection/deprotection procedure as described in Examples 24-26. The synthesis of thymine monomer is described in Examples 27-28, and that of the protected cytosine monomer is described in Example 29.

The synthesis of the protected adenine monomer (Figure 14) involved alkylation with ethyl bromoacetate (Example 30) and verification of the position of substitution by X-ray crystallography, as being the wanted 9-position. The No-amino group then was protected with the benzyloxycarbonyl group by 35 the use of the reagent N-ethyl-benzyloxycarbonylimidazole tetraflu r borate (Example 31). Simpl hydr lysis of the product ester (Example 32) gave No-benzyloxycarbonyl-9carboxym thyl adenine, which then was used in the standard procedure (Examples 33-34, Figur 13). The adenine monomer has been built into tw different PNA-oligomers (Examples 56, 57, 71 and 73).

The synthesis of the protected G-monomer is outlined in Figure 15. The starting material, 2-amino-6-chloropurine, was alkylated with bromoacetic acid (Example 35) and the chlorine atom was then substituted with a benzyloxy group (Example 36). The resulting acid was coupled to the (bocaminoethyl) glycine methyl ester (from Example 26) with agent PyBrop^m, and the resulting ester was hydrolysed (Example 37). The O⁶-benzyl group was removed in the final HF-cleavag step in the synthesis of the PNA-oligomer. Cleavage was verified by finding the expected mass of the final PNA-oligomer, upon incorporation into an PNA-oligomer using disopropyl carbodiimide as the condensation agent (Examples 55 and 71).

Extended Backbones

Alterations of the groups A, C and D (figure 16) is 20 demonstrated by the synthesis of monomeric building blocks and incorporation into PNA-oligomers.

In one example, the C group was a CH(CH₃) group. The synthesis of the corresponding monomer is outlined in Figure 17. It involves preparation of Boc-protected 1-amino-2,3-propanediol (Example 38), which is cleaved by periodate t give bocaminoacetaldehyde, which is used directly in the n xt reaction. The bocaminoacetaldehyde can be condensed with a variety of amines; in Example 39, alanine ethyl ester was used. In Examples 40-42, the corresponding thymine monomers were prepared. The monomer has been incorporated into an 8-mer (Example 60) by the DCC-coupling protocol (Examples 56 and 57).

In another example, the D group is a (CH₂)₃ group. The synthesis of the corresponding monomer is outlined in figure 35 18.A and described in Examples 43-44.

In another example, the A group is a $(CH_2)_2CO$ gr up. The synthesis f the corresponding thymine mon mer is outlined figure 18.B and Examples 46 through 48.

In yet another example, the C group is a (CH₂)₂ group.

5 The synthesis of the thymine and protected cytosine monomer is outlined in Figure 19 and Examples 49 through 54. Hybridization experiments with a PNA-oligomer containing ne unit is described in Examples 61 and 81, which shows a significant lowering of affinity but a retention of specificity.

General Remarks

The following abbreviations are used in the experimental examples: DMF, N,N-dimethylformamide; DCC, N,N-dicyclohexyl carbodiimide; DCU, N,N-dicyclohexyl urea; THF, tetrahydrofuran; aeg, N-acetyl (2'-aminoethyl)glycine; pfp, pentafluorophenyl; Boc, tert-butoxycarbonyl; Z, benzyloxy-carbonyl; NMR, nuclear magnetic resonance; s, singlet; d, doublet; dd, doublet of doublets; t; triplet; q, quartet; m, multiplet; b, broad; δ, chemical shift;

20 NMR spectra were recorded on either a JEOL FX 900 spectrometer, or a Bruker 250 MHz with tetramethylsilane as Mass spectrometry was performed on a internal standard. MassLab VG 12-250 quadropole instrument fitted with a VG FAB Melting points were recorded on Buchi source and probe. 25 melting point apparatus and are uncorrected. Dimethylformamide was dried over 4 Å molecular sieves. distilled and stored over 4 Å molecular sieves. Pyridine (HPLC quality) was dried and stored over 4 Å molecular sieves. Other solvents used were either the highest quality obtainable 30 or were distilled before use. Dioxane was passed through - basic alumina prior to use. Bocanhydride, 4-nitrophenol, bromoacetate, benzyloxycarbonyl methyl chloride, pentafluorophenol were all obtained through Aldrich Chemical Company. Thymine, cytosine, adenine were all obtained through 35 Sigma.

Thin layer chromatography (Tlc) was performed using the following s lvent systems: (1) chloroform:triethyl

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amine:methan 1, 7:1:2; (2) methylene chlorid :methanol, 9:1; (3) chloroform:methanol:ac tic acid 85:10:5. Spots were visualized by UV (254 nm) or/and spraying with a ninhydrin solution (3 g ninhydrin in 1000 ml 1-butanol and 30 ml acetic acid), after heating at 120°C for 5 min and, after spraying, heating again.

EXAMPLE 1

tert-Butyl 4-nitrophenyl carbonate

Sodium carbonate (29.14 g; 0.275 mol) and 4-nitrophen 1 (12.75 g; 91.6 mmol) were mixed with dioxane (250 ml). Bocanhydride (20.0 g; 91.6 mmol) was transferred to the mixtur with dioxane (50 ml). The mixture was refluxed for 1 h, cooled to 0°C, filtered and concentrated to 1/3, and then poured into water (350 ml) at 0°C. After stirring for 1/2 h., the product was collected by filtration, washed with water, and then dried over sicapent, in vacuo. Yield 21.3 g (97%). M.p. 73.0-74.5°C (litt. 78.5-79.5°C). Anal. for C11H13NO5 found(calc.) C: 55.20(55.23) H: 5.61(5.48) N: 5.82(5.85).

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EXAMPLE 2

(N'-Boc-2'-aminoethyl)glycine (2)

The title compound was prepared by a modification of the procedure by Heimer, et al. Int. J. Pept., 1984, 23, 203-25 211 N-(2-Aminoethyl)glycine (1, 3.00 g; 25.4 mmol) was dissolved in water (50 ml), dioxane (50 ml) was added, and the pH was adjusted to 11.2 with 2 N sodium hydroxide. trt-Butyl-4-nitrophenyl carbonate (7.29 g; 30.5 mmol) was dissolved in dioxane (40 ml) and added dropwise over a peri d of 2 h, during which time the pH was maintained at 11.2 with 2 N sodium hydroxide. The pH was adjusted periodically t 11.2 for three more hours and then the solution was left overnight. The solution was cooled to 0°C and the pH was carefully adjusted to 3.5 with 0.5 M hydrochloric acid. The 35 aqueous solution was washed with chloroform (3 x 200 ml), the pH adjust d t 9.5 with 2N sodium hydr xide and the solution was evaporat d to dryness, in vacuo (14 mmHg). The residue

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was extracted with DMF (25+2x10 ml) and th extracts filtered to remove excess salt. This results in a solution of the title compound in about 60% yield and greater than 95% purity by tlc (system 1 and visualised with ninhydrin, Rf=0.3). The 5 solution was used in the following preparations of Boc-aeg derivates without further purification.

EXAMPLE 3

N-1-Carboxymethylthymine (4)

This procedure is different from the literature 10 synthesis, but is easier, gives higher yields, and leaves no unreacted thymine in the product. To a suspension of thymine (3, 40.0 g; 0.317 mol) and potassium carbonate (87.7 g; 0.634 mmol) in DMF (900 ml) was added methyl bromoacetate (30.00 ml; 15 0.317 mmol). The mixture was stirred vigorously overnight under nitrogen. The mixure was filtered and evaporated to dryness, in vacuo. The solid residue was treated with water (300 ml) and 4 N hydrochloric acid (12 ml), stirred for 15 min at 0°C, filtered, and washed with water (2 x 75 ml). 20 precipitate was treated with water (120 ml) and 2N sodium hydroxide (60 ml), and was boiled for 10 minutes. The mixture was cooled to 0°C, filtered, and the pure title compound was precipitated by the addition of 4 N hydrochloric acid (70 ml). Yield after drying, in vacuo over sicapent: 37.1 g (64%). H- $DMSO-d_{\ell}):$ 25 NMR: (90 MHz: 11.33 ppm 7.49(d,J=0.92Hz,1H,ArH); 4.38 (s,2H,CH₂); 1.76 (d,J=0.92Hz,T- CH_3)

EXAMPLE 4

30 N-1-Carboxymethylthymine pentafluorophenyl ester (5)

N-1-Carboxymethylthymine (4, 10.0g; 54.3 mmol) and pentafluorophenol (10.0 g; 54.3 mmol) were dissolved in DMF (100 ml) and cooled to 5°C in ice water. DCC (13.45 g; 65.2 mmol) then was added. When the temperature passed below 5°C, 35 the ice bath was removed and the mixture was stirred for 3 h at ambient temperatur. The precipitated DCU was removed by filtration and washed twice with DMF (2 x 10 ml). The

combined filtrate was poured int ther (1400 ml) and cooled to 0°C. Petroleum ether (1400 ml) was added and th mixtur was left overnight. The title c mpound was is lated by filtration and was washed thoroughly with petroleum ether. 5 Yield: 14.8 g(78%). The product was pure enough to carry ut the next reaction, but an analytical sample was obtained by recrystallization from 2-propanol. M.p. 200.5-206°C Anal. f r C₁₃H₇F₅N₂O₄. Found(calc.) C: 44.79(44.59); H: 2.14(2.01) N: 8.13(8.00). FAB-MS: 443 (M+1+glycerol), 351 (M+1). H-NMR (90 MHz; DMSO-d₆): 11.52 ppm (s,1H,NH); 7.64 (s,1H,ArH); 4.99 (s,2H,CH₂); 1.76 (s,3H,CH₃).

EXAMPLE 5

1-(Boc-aeg) thymine (6)

To the DMF-solution from above was added triethyl amin 15 (7.08 ml; 50.8 mmol) followed by N-1-carboxymethylthymine pentafluorophenyl ester (5, 4.45 g; 12.7 mmol). The resultant solution was stirred for 1 h. The solution was cooled to 0°C and treated with cation exchange material ("Dowex 50W X-8", 20 40 g) for 20 min. The cation exchange material was remov d by filtration, washed with dichloromethane (2 x 15 ml), and dichloromethane (150 ml) was added. The resulting soluti n was washed with saturated sodium chloride, dried over magnesium sulfate, and evaporated to dryness, in vacuo, first 25 by a water aspirator and then by an oil pump. The residue was shaken with water (50 ml) and evaporated to dryness. procedure was repeated once. The residue then was dissolv d in methanol (75 ml) and poured into ether (600 ml) and petroleum ether (1.4 L). After stirring overnight, the white 30 solid was isolated by filtration and was washed with petroleum ether. Drying over sicapent, in vacuo, gave 3.50 g (71.7%). for C₁₆H₂₄N₄O₇. Found(calc.) M.p. 142-147°C. Anal. 49.59(50.00) H: 6.34(6.29) N: 14.58(14.58). H-NMR (250 MHz, DMSO-d_i): Due to the limited rotation around the secondary 35 amide bond several of the signals were doubled in the rati 2:1, (indicated in the list by mj. fr maj r and mi. fr minor). 12.73 ppm (b,1H, -CO₂H); 11.27 ppm (s, mj., imide);

11.25 ppm (s, mi., imide); 7.30 ppm (s, mj., ArH); 7.26 ppm (s, mi., ArH); 6.92 ppm (unres. t, mj., BocNH); 6.73 ppm (unres. t; mi., BocNH); 4.64 ppm (s, mj., T-CH₂-CO-); 4.47 ppm (s, mi., T-CH₂-CO-); 4.19 ppm (s, mi., CONRCH₂CO₂H); 3.97 ppm 5 (s, mj., CONRCH₂CO₂H); 3.41-2.89 ppm (unres. m, -CH₂CH₂- and water); 1.75 ppm (s, 3H, T-CH₃); 1.38 ppm (s, 9H, t-Bu). ¹³C-NMR: 170.68 ppm (CO); 170.34 (CO); 167.47 (CO); 167.08 (CO); 164.29 (CO); 150.9 (C5''); 141.92 (C6''); 108.04 (C2'); 77.95 and 77.68 (Thy-CH₂CO); 48.96, 47.45 and 46.70 (-CH₂CH₂- and NCH₂CO₂H); 37.98 (Thy-CH₃); 28.07 (t-Bu). FAB-MS: 407 (M+Na⁺); 385 (M+H⁺).

EXAMPLE 6

1-(Boc-aeg) thymine pentafluorophenyl ester (7, Boc-Taeg.OPfp) 15 1-(Boc-aeg) thymine (6) (2.00 g; 5.20 mmol) = was dissolved in DMF (5 ml) and methylene chloride (15 ml)-was added. Pentafluorophenol (1.05 g; 5.72 mmol) was added and the solution was cooled to 0°C in an ice bath. DDC then was added (1.29 g; 6.24 mmol) and the ice bath was removed after 20 2 min. After 3 h with stirring at ambient temperature, the precipitated DCU was removed by filtration and washed with methylene chloride. The combined filtrate was washed twice with aqueous sodium hydrogen carbonate and once with saturated sodium chloride, dried over magnesium sulfate, and evaporated 25 to dryness, in vacuo. The solid residue was dissolved in dioxane (150 ml) and poured into water (200 ml) at 0°C. The title compound was isolated by filtration, washed with water, and dried over sicapent, in vacuo. Yield: 2.20 g (77%). An analytical sample was obtained by recrystallisation from 2-30 propanol. M.p. 174-175.5°C. Analysis for C₂,H₂,N₄O₂F₅, found(calc.): C: 48.22(48.01); H: 4.64(4.21); N: 9.67(10.18). 'H-NMR (250 MHz, CDCl₁): Due to the limited rotation around the secondary amide bond several of the signals were doubled in the ratio 6:1 (indicated in the list by mj. for major and mi. 35 for minor). 7.01 ppm (s, mi., ArH); 6.99 ppm (s, mj., ArH); 5.27 ppm (unres. t, BocNH); 4.67 ppm (s, mj., T-CH,-CO-); 4.60 ppm (s, mi., T-CH₂-CO-); 4.45 ppm (s, mj., CONRCH₂CO₂Pfp); 4.42

ppm (s, mi., CONRCH₂CO₂Pfp); 3.64 ppm (t,2H,B cNHCH₂CH₂-); 3.87 ppm ("q",2H,B cNHCH₂CH₂-); 1.44(s,9H,t-Bu). FAB-MS: 551 (10; M+1); 495 (10; M+1-tBu); 451 (80; -B c).

5 EXAMPLE 7

N⁴-Benzyloxycarbonyl cytosine (9)

Over a period of about 1 h, benzyloxycarbonyl chloride (52 ml; 0.36 mol) was added dropwise to a suspension of cytosine (8, 20.0 g; 0.18 mol) in dry pyridine (1000 ml) at 0°C under nitrogen in oven-dried equipment. The solution then was stirred overnight, after which the pyridine suspension was evaporated to dryness, in vacuo. Water (200 ml) and 4 N hydrochloric acid were added to reach pH -1. The resulting white precipitate was filtered off, washed with water and partially dried by air suction. The still-wet precipitate was boiled with absolute ethanol (500 ml) for 10 min, cooled t 0°C, filtered, washed thoroughly with ether, and dried, in vacuo. Yield 24.7 g (54%). M.p.>250°C. Anal. for C12H11N3O3. Found(calc.); C: 58.59(58.77); H: 4.55(4.52); N: 17.17(17.13).

EXAMPLE 8 N'-Benzyloxycarbonyl-N'-carboxymethyl cytosine (10)

In a three necked round bottomed flask equipped with mechanical stirring and nitrogen coverage was placed methyl bromacetate (7.82 ml;82.6 mmol) and a suspension of N⁴-benzyloxycarbonyl-cytosine (9, 21.0 g;82.6 mmol) and potassium carbonate (11.4 g;82.6 mmol) in dry DMF (900 ml). The mixture was stirred vigorously overnight, filtered, and evaporated to dryness, in vacuo. Water (300 ml) and 4 N hydrochloric acid (10 ml) were added, the mixture was stirred for 15 minutes at 0°C, filtered, and washed with water (2 x 75 ml). The isolated precipitate was treated with water (120 ml), 2N sodium hydroxide (60 ml), stirred for 30 min, filter d, cool d to 0°C, and 4 N hydrochloric acid (35 ml) was add d. The title comp und was is lat d by filtrati n, wash d

thoroughly with water, recrystallized from methanol (1000 ml) and washed thoroughly with ether. This afford d 7.70 g (31%) of pure compound. The mother liquor from the recrystallization was reduced to a volume of 200 ml and cooled to 0°C.

This afforded an additional 2.30 g of a material that was purely by tlc but had a reddish color. M.p. 266-274°C. Anal. fr C₁₄H₁₃N₃O₅. Found(calc.); C: 55.41(55.45); H: 4.23(4.32); N: 14.04(13.86). H-NMR (90 MHz; DMSO-d₆): 8.02 ppm (d,J=7.32Hz,1H,H-6); 7.39 (s,5H,Ph); 7.01 (d,J=7.32Hz,1H,H-5); 5.19 (s,2H,PhCH₂-); 4.52 (s,2H).

EXAMPLE 9

N⁴-Benzyloxycarbonyl-N¹-carboxymethyl-cytosine pentafluorophenyl ester (11)

N'-Benzyloxycarbonyl-N'-carboxymethyl-cytosine (10, 15 4.00 g; 13.2 mmol) and pentafluorophenol (2.67 g; 14.5 mmol) were mixed with DMF (70 ml), cooled to 0°C with ice-water, and DCC (3.27 g; 15.8 mmol) was added. The ice bath was removed after 3 min and the mixture was stirred for 3 h at r om 20 temperature. The precipitated DCU was removed by filtration, washed with DMF, and the filtrate was evaporated to dryness, in vacuo (0.2 mmHg). The solid residue was treated with methylene chloride (250 ml), stirred vigorously for 15 min, filtered, washed twice with diluted sodium hydrogen carbonate 25 and once with saturated sodium chloride, dried over magnesium sulfate, and evaporated to dryness, in vacuo. residue was recrystallized from 2-propanol (150 ml) and the crystals were washed thoroughly with ether. Yield 3.40 g (55%). M.p. 241-245°C. Anal. for $C_{20}H_{12}N_3F_5O_5$. Found(calc.); 30 C: 51.56(51.18); H: 2.77(2.58); N: 9.24(8.95). H-NMR (90 MHz; CDCl₃): 7.66 ppm (d,J=7.63Hz,1H,H-6); 7.37 (s,5H,Ph); 7.31 (d, J=7.63Hz, 1H, H-5); 5.21 (s, 2H, PhCH₂-); 4.97 (s, 2H, NCH₂-).FAB-MS: 470 (M+1)

EXAMPLE 10

N'-Benzyl xycarb nyl-1-B c-a g-cyt sin (12)

To a solution of (N-Boc-2-amin ethyl)glycine (2) in DMF, prepared as described above, was added triethyl amine 5 (7.00 50.8 mmol) and N'-benzyloxycarbonyl-N'carboxymethyl-cytosine pentafluorophenyl ester (11, 2.70 g; 5.75 mmol). After stirring the solution for 1 h at room temperature, methylene chloride (150 ml), saturated sodium chloride (250 ml), and 4 N hydrochloric acid to pH -1 wer 10 added. The organic layer was separated and washed twice with saturated sodium chloride, dried over magnesium sulfate, and evaporated to dryness, in vacuo, first with a water aspirator and then with an oil pump. The oily residue was treated with water (25 ml) and was again evaporated to dryness, in vacuo. 15 This procedure them was repeated. The oily residue (2.80 g) was then dissolved in methylene chloride (100 ml), petrol um ether (250 ml) was added, and the mixture was stirr d overnight. The title compound was isolated by filtration and washed with petroleum ether. Tlc (system 1) indicated 20 substantial quantities of pentafluorophenol, but no attempt was made to remove it. Yield: 1.72 g (59%). 156°C(decomp.). H-NMR (250 MHz, CDCl₃): Due to the limited rotation around the secondary amide bond several of the signals were doubled in the ratio 2:1, (indicated in the list 25 by mj. for major and mi. for minor). 7.88 ppm (dd,1H,H-6); 7.39 (m,5H,Ph); 7.00 (dd,1H,H-5); 6.92 (b,1H,BocNH); 6.74 (b,1H,ZNH)-?; 5.19 (s,2H,Ph-CH3); 4.81 ppm (s, mj., Cyt-CH,-CO-); 4.62 ppm (s, mi., Cyt-CH,-CO-); 4.23 (s, mi., . CONRCH,CO,H); 3.98 ppm (s, mj., CONRCH,CO,H); 3.42-3.02 (unres. 30 m, -CH₂CH₂- and water);1.37 (s,9H,tBu). FAB-MS: 504 (M+1); 448 (M+1-tBu).

EXAMPLE 11

N⁴-Benzyloxycarbonyl-1-Boc-aeg-cytosine pentafluorophenyl 35 ester (13)

 N^4 -Benzyloxycarbonyl-1-Boc-aeg-cytosine (12, 1.50 g; 2.98 mmol) and pentafluorophenol (548 mg; 2.98 mmol) was

dissolved in DMF (10 ml) Methylene chl rid (10 ml) was added, the reaction mixture was cooled to 0°C in an ice bath, and DCC (676 mg; 3.28 mmol) was added. The ice bath was removed after 3 min and the mixture was stirred for 3 h at ambient 5 temperature. The precipitate was isolated by filtration and washed once with methylene chloride. The precipitate was dissolved in boiling dioxane (150 ml) and the solution was cooled to 15°C, whereby DCU precipitated. The DCU was removed by filtration and the resulting filtrate was poured into water 10 (250 ml) at 0°C. The title compound was isolated by filtration, was washed with water, and dried over sicapent, in vacuo. Yield 1.30 g (65%). Analysis for $C_{29}H_{28}N_5O_8F_5$. Found(calc.); C: 52.63(52.02); H: 4.41(4.22); N: 10.55(10.46). H-NMR (250 MHz; DMSO-d_k): showed essentially the spectrum of 15 the above acid, most probably due to hydrolysis of the ester. FAB-MS: 670 (M+1); 614 (M+1-tBu)

EXAMPLE 12

4-Chlorocarboxy-9-chloroacridine

4-Carboxyacridone (6.25 g; 26.1 mmol), thionyl chloride (25 ml), and 4 drops af DMF were heated gently under a flow of nitrogen until all solid material had dissolved. The solution then was refluxed for 40 min. The solution was cooled and excess thionyl chloride was removed in vacuo. The last traces of thionyl chloride were removed by coevaporation with dry benzene (dried over Na-Pb) twice. The remaining yellow powder was used directly in the next reaction.

EXAMPLE 13

30 4-(5-Methoxycarbonylpentylamidocarbonyl)-9-chloroacridine

Methyl 6-aminohexanoate hydrochloride (4.70 g; 25.9 mmol) was dissolved in methylene chloride (90 ml), cooled to 0°C, triethyl amine (15 ml) was added, and the resulting solution then was immediately added to the acid chloride from above. The roundbottomed flask containingthe acid chloride was cooled to 0°C in an ice bath. The mixture was stirred vigorously for 30 min at 0°C and 3 h at room temperature. The

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resulting mixtur was filter d to remove th remaining solids, which were washed with methylene chloride (20 ml). The redbrown methylene chloride filtrate was subsequently washed twice with saturated sodium hydrogen carbonate, once with 5 saturated sodium chloride, dried over magnesium sulfate, and evaporated to dryness, in vacuo. To the resulting oily substance was added dry benzene (35 ml) and ligroin (60-80°C, dried over Na-Pb). The mixture was heated to reflux. Activated carbon and celite were added and mixture was 10 refluxed for 3 min. After filtration, the title compound crystallised upon cooling with magnetic stirring. isolated by filtration and washed with petroleum ether. The product was stored over solid potassium hydroxide. Yield 5.0 g (50%).

15

EXAMPLE 14

4-(5-Methoxycarbonylpentyl) amidocarbonyl-9-[6'-(4''-nitrobenzamido) hexylamino]-aminoacridine

4-(5-Methoxycarbonylpentylamidocarbonyl)-9-20 chloroacridine (1.30 g; 3.38 mmol) and phenol (5 g) were heated to 80°C for 30 min under a flow of nitrogen, aft r which 6-(4'-nitrobenzamido)-1-hexylamine (897 mg; 3.38 mm 1) The temperature raised to 120°C for 2 h. reaction mixture was cooled and methylene chloride (80 ml) was 25 added. The resulting solution was washed three times with 2N sodium hydroxide (60 ml portions) and once with water, dried over magnesium sulfate, and evaporated to dryness, in vacuo. The resulting red oil (1.8 g) was dissolved in methylene chloride (40 ml), cooled to 0°C. Ether (120 ml) was added and 30 the resultant solution was stirred overnight. This results in a mixture of solid material and an oil. The solid was The solid and the oil were reisolated by filtration. dissolved in methylene chloride (80 ml) and added dropwise to cold ether (150 ml). After 20 minutes of stirring, the title 35 compound was isolated by filtration in the form of orange crystals. The pr duct was washed with ther and dri d in WO 92/20702 PCT/EP92/01219

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vacuo over potassium hydr xide. Yield 1.60 g (77%). M.p. 145-147°C.

EXAMPLE 15

5 4-(5-Carboxypentyl) amidocarbonyl-9-[6'-(4''-nitrobenzamido) hexylamino]-aminoacridine

4-(5-Methoxycarbonylpentyl)amidocarbonyl-9-[6'-(4''nitrobenzamido)hexylamino]aminoacridine (503 mg; 0.82 mmol)
was dissolved in DMF (30 ml), and 2 N sodium hydroxide (30 ml)

10 was added. After stirring for 15 min, 2 N hydrochloric acid
(35 ml) and water (50 ml) were added at 0°C. After stirring
for 30 min, the solution was decanted, leaving an oily
substance which was dissolved in boiling methanol (150 ml),
filtered and concentrated to 1/3 volume. To the methanol
15 solution were added ether (125 ml) and 5-6 drops of HCl in
ethanol. The solution was decanted after 1 h of stirring at
0°C. The oily substance was redissolved in methanol (25 ml)
and precipitated with ether (150 ml). The title compound was
isolated as yellow crystals after stirring overnight. Yield
20 417 mg (80%). M.p. 173°C (decomp.).

EXAMPLE 16

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(a) 4-(5-pentafluorophenyloxycarbonylpentyl) amidocarbonyl-9-[6'-(4''-nitrobenzamido) hexylamino]-aminoacridine(Acr¹Opfp)

The acid from above (300 mg; 0.480 mmol) was dissolved in DMF (2 ml) and methylene chloride (8 ml) was added. Pentafluorophenol (97 mg; 0.53 mmol), transferred with 2 x 2 ml of the methylene chloride, was added. The resulting solution was cooled to 0°C after which DCC (124 mg; 0.60 mmol) was subsequently added. The ice bath was removed after 5 minutes and the mixture was left with stirring overnight. The precipitated DCU was removed by centrifugation and the centrifugate was evaporated to dryness, in vacuo, first by a 35 water aspirator and then by an oil pump. The residu was dissolved in methylene chlorid (20 ml), filtered, and evaporated to dryness, in vacuo. The residue was again dissolved

in methyl n hl ride and petr leum ether (150 ml). A 1 ml portion of 5M HCl in ther was added. The solvent was r moved by decanting after 30 min of stirring at 0°C. The residual city substance was dissolved in methylene chloride (100 ml). Petroleum ether (150 ml) was added and the mixture was left with stirring overnight. The next day the yellow precipitat d crystalline material was isolated by filtration and was washed with copious amounts of petroleum ether. Yield, after drying, 300 mg (78%). M.p. 97.5°C (decomp.) All samples show d satisfactory elemental analysis, H- and C-NMR and mass spectra.

(b) Experimental for the synthesis of PNA compounds, of. Figure 8

Materials: Boc-Lys (C1Z), benzhydrylamine-cop ly15 (styrene-1t-divinylbenzene) resin (BHA resin), and pmethylbenzhydrylamine-copoly(styrene-1t-divinylbenzene)r sin
(MBHA resin) were purchased from Peninsula Laboratori s.
Other reagents and solvents were: Biograde trifluoroacetic
acid from Halocarbon Products; diisopropylethylamine (99%; was
20 not further distilled) and N-acetylimidazole (98%) fr m
Aldrich: H₂O was distilled twice; anhydrous HF from Union
Carbide; synthesis grade N,N-dimethylformamide and analytical
grade methylene chloride (was not further distilled) fr m
Merck; HPLC grade acetonitrile from Lab-Scan; purum grade
25 anisole, N,N'-dicyclohexylcarbodiimid,
diisopropylcarbodiimide, puriss. grade 2,2,2-trifluoroethanol
from Fluka and trifluoromethanesulfonic acid from flourad.

(b) General Methods and Remarks

Except where otherwise stated, the following applies.

30 The PNA compounds were synthezised by the stepwise solid-phase approach (Merrifield, J. Am. Chem. Soc., 1963, 85, 2149) employing conventional peptide chemistry utilizing the TFA-labile tert-butyloxycarbonyl (Boc) group for "temporary" N-labile tert-butyloxycarbonyl (Boc) group for "temporary" N-protection (Merrifield, J. Am. Chem. Soc., 1964, 86, 304) and protection (Merrifield, J. Am. Chem. Soc., 1964, 86, 304) and chlorob nzyl xycarbonyl (Cl2) groups for "permanent" side chain prot ction. To btain C-t rminal amides, the PNAs were assembly do note the HF-labile BHA or MBHA resins (the MBHA).

resin has increased susceptibility to the final HF cleavage relative to the unsubstituted BHA resin (Matsueda, et al., Peptides, 1981, 2, 45). All reactions (except HF reactions) were carried out in manually operated standard solid-phase reaction vessels fitted with a coarse glass frit (Merrifield, et al., Biochemistry, 1982, 21, 5020). The quantitative ninhydrin reaction (Kaiser test), originally developed by Merrifield and co-workers (Sarin, et al., Anal. Biochem., 1981, 117, 147) for peptides containing "normal" amino acids, was successfully appplied (see Table I - III) using the "normally" employed effective extinction coefficient \(\epsilon = 15000 \) M'cm' for all residues to determine the completeness of the individual couplings as well as to measure the number of growing peptide chains. The theoretical substitution

15 S_{n-1} upon coupling of residue number n (assuming both complete deprotection and coupling as well as neither chain termination nor loss of PNA chains during the synthetic cycle) is calculated from the equation:

 $S_n = S_{n-1} \times (1 + (S_{n-1} \times \Delta MW \times 10^{-3} \text{ mmol/mol}))^{-1}$

20 where ΔMW is the gain in molecular weight ([ΔMW] = g/mol) and S., is the theoretical substitution upon coupling of the preceding residue n-1 ([S] = mmol/g). The estimated value (%) on the extent of an individual coupling is calculated relative to the measured substitution (unless S was not determined) and 25 include correction for the number of remaining free amino groups following the previous cycle. HF reactions were carried out in a Diaflon HF apparatus from Toho Kasei (Osaka, Japan). Vydac C_{18} (5 μ m, 0.46 x 25 cm and 5 μ m, 1.0 x 25 cm) reverse-phase columns, respectively were used for analytical 30 and semi-preparative HPLC on an SP8000 instrument. Buffer A was 5 vol % acetonitrile in water containing 445 μl trifluoroacetic acid per liter, and buffer B was 60 vol % acetonitrile in water containing 390 µl trifluoroacetic acid per liter. The linear gradient was 0-100% of buffer B in 30 35 min, flow rates 1.2 ml/min (analytical) and 5 ml/min (semi-The eluents w re monitored at 215 nm (analytical) and 230 nm (semi-preparative). Molecular weights

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f th PNAs were determin d by ²⁵²Cf plasma desorption time-of-flight mass spectrom try from the m an f the most abundant isotopes.

- 5 EXAMPLE 17
 Solid-Phase Synthesis of Acr1-[Taeg]₁₅-NH₂ and Shorter
 Derivatives
 - Derivatives

 (a) Stepwise Assembly of Boc-[Taeg]₁₅-BHA Resin

 The synthesis was initiated on 100 mg of preswollen and
- 10 neutralized BHA resin (determined by the quantitative ninhydrin reaction to contain 0.57 mmol NH₂/g) employing single couplings ("Synthetic Protocol 1") using 3.2 equivalents of BocTaeg-OPfp in about 33% DMF/CH₂Cl₂. The individual coupling reactions were carried out by shaking for
 - 15 at least 12 h in a manually operated 6 ml standard solid-phas reaction vessel and unreacted amino groups were blocked by acetylation at selected stages of the synthesis. The progr ss of chain elongation was monitored at several stages by th quantitative ninhydrin reaction (see Table I). Portions f
 - 20 protected Boc-[Taeg]₅-BHA, Boc-[Taeg]₁₀-BHA, and Boc-[Taeg]₁₅-BHA resins were taken out after assembling 5, 10, and 15 residues, respectively.

000

30

	Synthetic Step	Residue Coupled	and the state of t		s After	Estimated Extent of Coupling	
			Measd	Theoretol .	Single Coupling	Acetylation	(%)
	•0•		0.57				
5	1	BocTaeg	ND	0.50	1.30		<99.7
	2	BocTaeg	ND	0.44	1.43		<99.9
	3	BocTaeg	0.29	0.39	3.33		99.3
	4	BocTaeg	0.27	0.35	13.30		96.3
	5	BocTaeg	0.26	0.32	8.33		>99.9
0	6	BocTaeg	ND	0.30	7.78		>99.9
l	7	BocaTeg	ND	0.28	13.81	7.22	<97.8
	8	BocTaeg	ND	0.26	14.00	3∗	· <99.9
	9	BocTaeg	ND	0.24	30.33		93.2
	10	BocTaeg	0.16	0.23	11.67	2.67	>99.9
5	11	BocTaeg	ND	0.21	4.58		>99.9
	12	BocTaeg	ND	0.20	5.87		<99.4
	13	BocTaeg	ND	0.19	1.67		>99.9
	14	BocTaeg	ND	0.18	14.02		<93.1
	15	BocTaeg	0.07	0.17	4.20	3.33	>99.9

(b) Synthesis of Acr 1-[Taeg] 15-BEA Resin

Following deprotection of the residual Boc-[Taeg]₁₅-BHA resin (estimated dry weight is about 30 mg; ~0.002 mmol growing chains), the H-[Taeg]₁₅-BHA resin was reacted with about 50 equivalents (80 mg; 0.11 mmol) of Acr¹-OPfp in 1 ml of about 66% DMF/CH₂Cl₂ (i.e., a 0.11 M solution of th pentafluorophenylester) in a 3 ml solid-phase reaction vessel. As judged by a qualitative ninhydrin reaction, coupling of the acridine moiety was close to quantitative.

(c) Cleavage, Purification, and Identification of H-[Taeg]₅-NH₂

A portion of protected Boc-[Taeg]₅-BHA resin was treated with 50% trifluoroacetic acid in methylene chloride to remove the N-terminal Boc group (which is a precursor of

the potentially harmful tert-butyl cati n) pri r to th HF cleavag . Foll wing n utralization and washing (perf rmed in a way similar t th se f steps 2-4 in "Synth tic Protocol 1"), and drying for 2 h in vacuum, the resulting 67.1 mg (dry 5 weight) of H-[Taeg],-BHA resin was cleaved with 5 ml Hr:anisole (9:1, v/v) stirring at 0°C for 60 min. . Aft r removal of HF, the residue was stirred with dry diethyl ether (4 x 15 ml, 15 min each) to remove anisole, filtered under gravity through a fritted glass funnel, and dried. The PNA 10 was then extracted into a 60 ml (4 x 15 ml, stirring 15 min each) 10% aqueous acetic acid solution. Aliquots of this solution were analyzed by analytical reverse-phase HPLC to establish the purity of the crude PNA. The main peak at 13.0 min accounted for about 93% of the total absorbance. The 15 remaining solution was frozen and lyophilized to afford about 22.9 mg of crude material. Finally, 19.0 mg of the crude product was purified from five batches, each containing 3.8 mg in 1 ml of H2O. The main peak was collected by use of a semi-preparative reverse-phase column. Acetonitrile was 20 removed on a speed vac and the residual solution was frozen (dry ice) and subsequently lyophilized to give 13.1 mg of >99% The PNA molecule readily dissolved in pure H-[Taeg],-NH,. water and had the correct molecular weight based on mass spectral determination. For (M+H) the calculated m/z value 25 was 1349.3 and the measured m/z value was 1347.8.

(d) Cleavage, Purification, and Identification of H[Taeg]₁₀-MH₂

A portion of protected Boc-[Taeg]₁₀-BHA resin was treated as described in section (c) to yield 11.0 mg of crud material upon HF cleavage of 18.9 mg dry H-[Taeg]₁₀-BHA resin. The main peak at 15.5 min accounted for about 53% of the t tal absorbance. About 1 mg of the crude product was purified repeatedly (for reasons described below) to give approximately 0.1 mg of at least 80% but presumably >99% pure H-[Taeg]₁₀-NH₂. A rather broad tail eluting after the target peak and accounting for about 20% of th t tal absorbanc c uld n t be rem ved (only slightly r duc d) upon th r peated

purification. Judged by the mass spectrum, which only confirms the presence of the correct molecular weight H[Taeg]₁₀-NH₂, the tail phenomonen is ascribed to more or less well-defined aggregational/conformational states of the target molecule. Therefore, the crude product is likely to contain more than the above-mentioned 53% of the target molecule. H[Taeg]₁₀-NH₂ is readily dissolved in water. For (M+H)* the calculated m/z value was 2679.6 and the measured m/z value was 2681.5.

10 (e) Cleavage, Purification, and Identification of H[Taeg]₁₅-NH₂.

A portion of protected Boc-[Taeg]₁₅-BHA resin was treated as described in section (c) to yield 3.2 mg of crud material upon HF cleavage of 13.9 mg dry H-[Taeg]₁₅-BHA resin.

15 The main peak at 22.6 min was located in a broad bulge accounting for about 60% of the total absorbance (Fig. 12a). Again (see the preceding section), this bulge is ascribed to aggregational/conformational states of the target molecul H-[Taeg]₁₅-NH2 since mass spectral analysis of the collected 20 "bulge" did not significantly reveal the presence of other molecules. All of the crude product was purified collecting the "bulge" to give approximately 2.8 mg material. For (M+Na) the calculated m/z value was 4033.9 and the measured m/z value was 4032.9.

25 (f) Cleavage, Purification, and Identification f $Acr^{1}-[Taeg]_{15}-NH_{2}.$

A portion of protected Acr¹-[Taeg]₁₅-BHA resin was treated as described in section (b) to yield 14.3 mg of crude material upon HF cleavage of 29.7 mg dry Acr¹-[Taeg]₁₅-BHA resin. Taken together, the main peak at 23.7 min and a "dimer" (see below) at 29.2 min accounted for about 40% of the total absorbance (Fig. 12b). The crude product was purified repeatedly to give approximately 1 mg of presumably >99% pure Acr¹-[Taeg]₁₅-NH₂ "contaminated" with self-aggregated molecules eluting at 27.4 min, 29.2 min, and finally as a huge broad bulg luting with 100% buffer B (Fig. 12c). This interpretation is in agree ment with the observation that those

peaks gr w upon standing (f r h urs) in aqu us acetic acid soluti n, and finally precipitate ut quantitativ ly. F r $(M+H)^+$ the calculat d m/z value was 4593.6 and the measured m/z value was 4588.7.

- (g) Synthetic Protocol 1
- (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 3 ml, 3 x 1 min and 1 x 30 min; (2) washing with CH,Cl,, 3 ml, 6 x 1 min; (3) neutralization with DIEA/CH,Cl, (1: 19, v/v), 3 ml, 3 x 2 min; (4) washing with CH,Cl,, 3 ml, 6 x 1 min, and drain 10 for 1 min; (5) 2-5 mg sample of PNA-resin may be taken out and dried thoroughly for a quantitative ninhydrin analysis t determine the substitution; (6) addition of 3.2 equiv. (0.18 mmol; 100 mg) BocTaeg-OPfp dissolved in 1 ml CH,Cl, foll w d by addition of 0.5 ml DMF (final concentration f 15 pentafluorophenylester ~0.12 M); the coupling reaction was allowed to proceed for a total of 12-24 h shaking at ro m temperature; (7) washing with DMF, 3 ml, 1 x 2 min; (8) washing with CH,Cl2, 3 ml, 4 x 1 min; (9) neutralization with DIEA/CH,Cl, (1: 19, V/V), 3 ml, 2 x 2 min; (10) washing with 20 CH,Cl,, 3 ml, 6 x 1 min; (11) 2-5 mg sample of protected PNAresin is taken out for a rapid qualitative ninhydrin test and further 2-5 mg is dried thoroughly for a quantitative ninhydrin analysis to determine the extent of coupling (after cycles 7, 10, and 15 unreacted amino groups were blocked by 25 acetylation with N-acetylimidazol in methylene chloride).

EXAMPLE 18

Solid-Phase Synthesis of Acr1-[Taeg] 15-Lys-NH2 and Sh rter Derivatives

30 (a) Stepwise Assembly of Boc-[Taeg]₁₅-Lys(Cl2)-BHA Resin

The synthesis was initiated by a quantitative loading (standard DCC in situ coupling in neat CH_2Cl_2) of Boc-Lys(Cl2) onto 100 mg of preswollen and neutralized BHA resin (0.57 mmol NH_2/g). Further extension of the protected PNA chain empl yed single couplings ("Synthetic Protocol 2") for cycl s 1 t 5 and cycles 10 to 15 using 3.2 equival nts f BocTa g-

OPfp in about 33% DMF/CH₂Cl₂. Cycles 5 to 10 employed an extra straight DCC (i.e., in situ) coupling of the free acid BocTaeg-OH in about 33% DMF/CH₂Cl₂. All coupling reactions were carried out by shaking for at least 12 h in a manually operated 6 ml standard solid-phase reaction vessel. Unreacted amino groups were blocked by acetylation at the same stages of the synthesis, as was done in Example 17. Portions of protected Boc-[Taeg]₅-Lys(Cl2)-BHA and Boc-[Taeg]₁₀-Lys(Cl2)-BHA resins were taken out after assembling 5 and 10 PNA residues, respectively. As judged by the analytical HPLC chromatogram of the crude cleavage product from the Boc-[Taeg]₁₀-Lys(Cl2)-BHA resin (see section (e)), an additional "free acid" coupling of PNA residues 5 to 10 gave no significant improvement of the synthetic yield as compared to the throughout single-coupled residues in Example 17.

- (b) Synthesis of Acr¹-[Taeg]₁₀-Lys(Cl2)-BHA Resin
 Following deprotection of a portion of Boc-[Taeg]₁₀Lys(Cl2)-BHA resin (estimated dry weight is about 90 mg;
 0.01 mmol growing chains), the H-[Taeg]₁₅-BHA resin was
 20 reacted with about 20 equivalents (141 mg; 0.19 mmol) of Acr¹OPfp in 1 ml of about 66% DMF/CH₂Cl₂ in a 3 ml solid-phase
 reaction vessel. As judged by a qualitative ninhydrin
 reaction, coupling of the acridine moiety was close to
 quantitative.
- (c) Synthesis of Acr¹-[Taeg]₁₅-Lys(Cl2)-BHA Resin
 Following deprotection of the residual Boc-[Taeg]₁₅Lys(Cl2)-BHA resin (estimated dry weight about 70 mg; ~ 0.005
 mmol growing chains), the H-[Taeg]₁₅-Lys(Cl2)-BHA resin was
 reacted with about 25 equivalents (91 mg; 0.12 mmol) of Acr¹OPfp in 1 ml of about 66% DMF/CH₂Cl₂ in a 3 ml solid-phase
 reaction vessel. As judged by a qualitative ninhydrin
 reaction, coupling of the acridine moiety was close to
 quantitative.
- (d) Cleavage, Purification, and Identification of H-35 [Taeg]₅-Lys-NH₂

A p rti n f pr tected B c-[Taeg]_{\S}-Lys(ClZ)-BHA resin was treated as described in Example 17c t yield 8.9 mg of

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crude material upon HF cl avage f 19.0 mg dry H-[Ta g]₅-Lys(Cl2)-BHA resin. The main peak at 12.2 min (eluted at 14.2 min if injected fr m an aqu us solution instead f the 10% aqueous acetic acid solution) accounted for about 90% of the total absorbance. About 2.2 mg of the crude product was purified to give approximately 1.5 mg of 99% pure H-[Taeg]₅-Lys-NH₂.

(e) Cleavage, Purification, and Identification of H-[Taeg] 10-Lys-NH2

A portion of protected Boc-[Taeg]₁₀-Lys(Cl2)-BHA resin was treated as described in Example 17c to yield 1.7 mg of crude material upon HF cleavage of 7.0 mg dry H-[Taeg]₁₀-Lys(Cl2)-BHA resin. The main peak at 15.1 min (eluted at 17.0 min if injected from an aqueous solution instead of the 10% aqueous acetic acid solution) accounted for about 50% of the total absorbance. About 1.2 mg of the crude product was purified to give approximately 0.2 mg of >95% pure H-[Taeg]₁₀-Lys-NH₂. Figure 13a. For (M+H) the calculated m/z value was 2807.8 and the measured m/z value was 2808.2.

20 (f) Cleavage, Purification, and Identification f
Acri-[Taeg];0-Lys-NH2

99.1 mg protected Acr - [Taeg] 10-Lys(Cl2)-BHA resin (dry weight) was cleaved as described in Example 17c to yield 42.2 mg of crude material. The main peak at 25.3 min (eluted at 25.3 min if injected from an aqueous solution instead of the 10% aqueous acetic acid solution) accounted for about 45% f the total absorbance. An 8.87 mg portion of the crude product was purified to give approximately 5.3 mg of >97% pure H-[Taeg] 10-Lys-NH2. For (M+H) the calculated m/z value was 2849.8.

(g) Cleavage and Purification of Acr¹-[Taeg]₁₅-Lys-NH₂
A 78.7 mg portion of protected Acr¹-[Taeg]₁₅-Lys(ClZ)BHA resin (dry weight) was cleaved as described in Example I section (c) to yield 34.8 mg of crude material. The main peak
35 at 23.5 min (about the same elution time if injected fr m an aque us solution instead of th 10% aqu us acetic acid s lution) and a "dimer" at 28.2 min acc unted f r about 35%

of the total absorbanc. About 4.5 mg of the crude product was purified to giv approximately 1.6 mg f presumably >95% pure H-[Taeg]₁₀-Lys-NH₂. This compound could not be free of the "dimer" peak, which grew upon standing in aqueous acetic acid solution.

(h) Synthetic Protocol 2

(1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 3 ml, 3 x 1 min and 1 x 30 min; (2) washing with CH_2Cl_2 , 3 ml, 6 x 1 min; (3) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 3 ml, 10 3 x 2 min; (4) washing with CH₂Cl₂, 3 ml, 6 x 1 min, and drain for 1 min; (5) 2-5 mg sample of PNA-resin can be taken out and dried thoroughly for a qualitative ninhydrin analysis; (6) for cycles 1 to 5 and cycles 10 to 15 the coupling reaction was carried out by addition of 3.2 equiv. (0.18 mmol; 100 mg) 15 BocTaeg-OPfp dissolved in 1 ml CH,Cl, followed by addition of 0.5 ml DMF (final concentration of pentafluorophenylester -0.12 M); the coupling reaction was allowed to proceed for a total of 12-24 h with shaking; cycles 5 to 10 employed an additional 0.12 M DCC coupling of 0.12 M BocTaeg-OH in 1.5 ml 20 DMF/CH₂Cl₂ (1:2, v/v); (7) washing with DMF, 3 ml, 1 x 2 min; (8) washing with CH_2Cl_2 , 3 ml, 4 x 1 min; (9) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 3 ml, 2 x 2 min; (10) washing with CH₂Cl₂, 3 ml, 6 x 1 min; (11) 2-5 mg sample of protected PNA-resin is taken out for a qualitative ninhydrin test (after 25 cycles 7, 10, and 15 unreacted amino groups were blocked by acetylation with N-acetylimidazol in methylene chloride).

EXAMPLE 19

Improved Solid-Phase Synthesis of H-[Taeg] 10-Lys-NH,

The protected PNA was assembled onto an MBHA resin, using approximately half the loading of the BHA resin used in the previous examples. Furthermore, all cycles except one was followed by acetylation of uncoupled amino groups. The following describes the synthesis in full detail:

(a) Preparati n f B c-Lys(ClZ)-NH-CH(p-CH₃-C₆H₄)-C₆H₄

Resin (MBHA R sin) with an Initial Substitution
f 0.3 mm 1/g

The desired substitution of Boc-Lys(ClZ)-MBHA resin was 5 0.25 - 0.30 mmol/g. In order to get this value, 1.5 mmol of Boc-Lys(ClZ) was coupled to 5.0 g of neutralized and preswollen MBHA resin (determined by the quantitativ ninhydrin reaction to contain 0.64 mmol NH2/g) using a singl "in situ" coupling (1.5 mmol of DCC) in 60 ml of CH2Cl2. Th 10 reaction was carried out by shaking for 3 h in a manually operated, 225 ml, standard, solid-phase reaction vess 1. Unreacted amino groups were then blocked by acetylation with a mixture of acetic anhydride/pyridine/CH2Cl2 (1:1:2, v/v/v) A quantitative ninhydrin reaction on th for 18 h. 15 neutralized resin showed that only 0.00093 mmol/g free amin remained (see Table I), i.e. 0.15% of the original amino The degree of substitution was estimated by deprotection and ninhydrin analysis, and was found to be 0.32 mmol/g for the neutralized H-Lys(Cl2)-MBHA resin. 20 compares well with the maximum value of 0.28 mmol/g for a quantitative coupling of 0.30 mmol Boc-Lys(ClZ)/g resin (see Table II).

- (b) Stepwise Assembly of Boc-[Taeg]₃-Lys(Cl2)-MBHA Resin
- The entire batch of H-Lys(ClZ)-MBHA resin prepared in section (a) was used directly (in the same reaction vess 1) to assemble Boc-[Taeg]₃-Lys(ClZ)-MBHA resin by singl couplings ("Synthetic Protocol 3") utilizing 2.5 equivalents of BocTaeg-OPfp in neat CH₂Cl₂. The quantitative ninhydrin reaction was appplied throughout the synthesis (see Table II).
 - (c) Stepwise Assembly of Boc-[Taeg]₈-Lys(Cl2)-MBHA
 Resin

About 4.5 g of wet Boc-[Taeg]₃-Lys(ClZ)-MBHA resin (-0.36 mmol growing chains; taken out of totally - 19 g wet 35 resin prepared in section (b)) was placed in a 55 ml SPPS raction v ssel. B-c-[Taeg]₈-Lys(ClZ)-MBHA resin was assembled by single uplings ("Synthetic Protocol 4")

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utilizing 2.5 equivalents of BocTaeg-OPfp in ab ut 30% DMF/CH_2Cl_2 . The progress of the synthesis was monitored at all stages by the quantitative ninhydrin reaction (see Table II).

(d) Stepwise Assembly of Boc-[Taeg]₁₀-Lys(ClZ)-MBHA Resin

About 1 g of wet Boc-[Taeg]₈-Lys(Cl2)-MBHA resin (-0.09 mmol growing chains; taken out of totally -4 g wet resin prepared in section (c)) was placed in a 20 ml SPPS reaction vessel. Boc-[Taeg]₁₀-Lys(Cl2)-MBHA resin was assembled by th single-coupling protocol employed in the preceding section utilizing 2.5 equivalents of BocTaeg-OPfp in about 30% DMF/CH₂Cl₂. The reaction volume was 3 ml (vigorous shaking). The synthesis was monitored by the quantitative ninhydrin reaction (see Table II).

·	Synthetic Step	Residue Coupled	Substitution After Deprotection (mmol/g)		Remaining Group (µm	Estimated Extent of Coupling	
			Measd	Theoret	Single Coupling	Acetylation	(%)
20	•0•	BocLys(CIZ)	0.32	0.28		0.93	
	1	BocTaeg	0.23	0.26	0.97	0.54	> 99.9
	2	BocTaeg	0.21	0.24	0.92	0.46	99.8
	3	BocTaeg	0.19	0.23	1.00	0.57	99.7
	4	BocTaeg	0.18	0.21	1.85		99.3
25	5	BocTaeg	0.17	0.20	2.01	0.19	99.9
	6	BocTaeg	0.15	0.19	1.69	0.10	99.0
	7	BocaTeg	0.11	0.18	1.11	0.66	99.1
	8	BocTaeg	0.12	0.17	1.82	0.44	99.0
	9	BocTaeg	0.10	0.17	5.63	0.56	94.8
30	10	BocTaeg	0.11	0.16	1.54	0.67	99.1

(e) Synthesis of Ac-[Taeg] 10-Lys(ClZ)-MBHA Resin

Following depr t ction f a p rti n f Boc-[Taeg]₁₀-Lys(ClZ)-MBRA resin (estimated dry weight is about 45 mg), the 35 resin was next ac tylated quantitatively with a 2 ml mixture

of acetic anhydride/pyridine/CH₂Cl₂ (1:1:2, v/v/v) f r 2 h in a 3 ml solid-phase r action vessel.

(f) Cleavage, Purificati n, and Identificati n f H[Taeg] 10-Lys-NH2

A portion of protected Boc-[Taeg] 10-Lys(ClZ)-BHA resin 5 was treated as described in Example 17c to yield about 24 mg of crude material upon HF cleavage of 76 mg dry H-[Taeg]5-Lys(C12)-BHA resin. The main peak at 15.2 min (which includes impurities such as deletion peptides and various byproducts) 10 accounted for about 78% of the total absorbance. peak also accounted for about 88% of the "main peak plus deletion peaks" absorbance, which is in good agreement with the overall estimated coupling yield of 90.1% obtained by summarizing the individual coupling yields in Table II. A 7.2 15 mg portion of the crude product was purified from two batches by use of a semi-preparative reserse-phase column, (collecting the main peak in a beaker cooled with dry ice/2-propan 1). Each contained 3.6 mg in 1 ml of H2O. The frozen solution was lyophilized directly (without prior removal of acetonitrile 20 on a speed vac) to give 4.2 mg of 82% pure H-[Taeg]₁₀-Lys-NH₂.

> (g) Cleavage, Purification, and Identification of Ac-[Taeg]₁₀-Lys-NH₂

A 400.0 mg portion of protected Ac-[Taeg]₁₀-Lys(ClZ)-BHA resin (dry weight) was cleaved as described in Example 17c, except for the TFA treatment to yield 11.9 mg of crude material. The main peak at 15.8 min accounted for about 75% of the total absorbance. A 4.8 mg portion of the crude product was purified to give approximately 3.5 mg of >95% pure Ac-[Taeg]₁₀-Lys-NH₂. For (M+H) the calculated m/z value = 2849.8 and the measured m/z value = 2848.8.

- (h) Synthetic Protocol 3.
- (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 100 ml, 3 x 1 min and 1 x 30 min; (2) washing with CH₂Cl₂, 100 ml, 6 x 1 min; (3) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 35 100 ml, 3 x 2 min; (4) washing with CH₂Cl₂, 100ml, 6 x 1 min, and drain f r 1 min; (5) 2-5 mg sample of PNA-resin is taken out and dri d thor ughly f r a quantitative ninhydrin analysis

t determine the substitution; (6) addition of 2.5 equiv. (3.75 mmol; 2.064 g) BocTaeg-OPfp dissolved in 35 ml CH,Cl, (final concentration of pentafluorophenylester -0.1 M); the coupling reaction was allowed to proceed for a total of 20-24 5 h with shaking; (7) washing with DMF, 100 ml, 1×2 min (to remove precipitate of BocTaeg-OH); (8) washing with CH2Cl2, 100 ml, 4 x 1 min; (9) neutralization with DIEA/CH,Cl, (1: 19, v/v), 100 ml, 2 x 2 min; (10) washing with CH_2Cl_2 , 100 ml, 6 x 1 min; (11) 2-5 mg sample of protected PNA-resin is taken 10 out for a rapid qualitative ninhydrin test and a further 2-5 mg is dried thoroughly for a quantitative ninhydrin analysis to determine the extent of coupling; (12) blocking of unreacted amino groups by acetylation with a 100 ml mixture of acetic anhydride / pyridine / CH2Cl2 (1:1:2, v/v/v) for 2 15 h; (13) washing with CH_2Cl_2 , 100 ml, 6 x 1 min; (14) 2 x. 2-5 mg samples of protected PNA-resin are taken out, neutralized with DIEA/CH,Cl2 (1: 19, v/v) and washed with CH2Cl2 for qualitative and quantitative ninhydrin analyses.

(i) Synthetic Protocol 4.

20 (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 25 ml, 3 x 1 min and 1 x 30 min; (2) washing with CH,Cl,, 25 ml, 6 x 1 min; (3) neutralization with DIEA/CH,Cl, (1: 19, v/v), 25 ml, 3 x 2 min; (4) washing with CH_2Cl_2 , 25 ml, 6 x 1 min, and drain for 1 min; (5) 2-5 mg sample of PNA-resin is taken ut 25 and dried thoroughly for a quantitative ninhydrin analysis t determine the substitution; (6) addition of 2.5 equiv. (0.92 mmol; 0.506 g) BocTaeg-OPfp dissolved in 6 ml CH,Cl, followed addition of 3 ml DMF (final concentration of pentafluorophenylester -0.1 M); the coupling reaction was 30 allowed to proceed for a total of 20-24 hrs with shaking; (7) washing with DMF, 25 ml, 1 x 2 min; (8) washing with CH,Cl,, 25 ml, 4 x 1 min; (9) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 25 ml, 2 x 2 min; (10) washing with CH_2Cl_2 , 25 ml, 6 x 1 min; (11) 2-5 mg sample of protected PNA-resin is taken out 35 for a rapid qualitativ ninhydrin test and a further 2-5 mg is dried ther ughly for a quantitative ninhydrin analysis to determine the extent of coupling; (12) blocking of unreacted amino groups by acetylation with a 25 ml mixtur of acetic anhydride/pyridine/CH₂Cl₂ (1:1:2, v/v/v) f r 2 h (except after the first cycle); (13) washing with CH₂Cl₂, 25 ml, 6 x 1 min; (14) 2 x 2-5 mg samples of protected PNA-resin are taken ut, neutralized with DIEA/CH₂Cl₂ (1: 19, v/v) and washed with CH₂Cl₂ for qualitative and quantitative ninhydrin analyses.

EXAMPLE 20 Solid-Phase Synthesis of H-[Taeg]₅-Caeg-[Taeg]₄-Lys-NH₂ 10 (a) : Stepwise Assembly of Boc-[Taeg]₅-C(z)aeg-[Taeg]₄Lys(Cl2)-MBHA Resin

About 2.5 g of wet Boc-[Taeg]₃-Lys(Cl2)-MBHA resin (-1/6 of the total remaining about 16 g wet resin; -0.75 g dry resin -0.15 mmol growing chains) was placed in a 6 ml SPPS reaction vessel. Boc-[Taeg]₅-Caeg-[Taeg]₄-Lys(Cl2)-MBHA'r sin was assembled by double coupling of all Taeg-residues utilizing the usual 2.5 equivalents of BocTaeg-OPfp in 2.5 ml about 30% DMF/CH₂Cl₂, except that the first residue was single-coupled. Incorporation of the C(Z)aeg-residue was accomplished by coupling with 2.0 equivalents of BocC(Z)a g-OPfp in TFE/CH₂Cl₂ (1:2, v/v). The progress of the synthesis was monitored at all stages by the quantitative ninhydrin reaction (see Table III).

25	Synthetic Step	Residue Coupled	Substitution After Deprotection (mmol/g)		Remaining Free Amino Groups After (umol/g)			Estimated Extent of Coupling
			Messd.	Theoret.	1st Coupl	2nd Coupl	Acetyl- stion	
İ	3		0.19	0.23	1.00		0.57	
İ	4	BocTaeg	0.17	0.21	4.88		97.3	97.3
30	5	BocC(Z)aeg	0.11	0.20	70.20	27.98	1.33	78.4 (46)
	6	BocTaeg	0.10	0.19	24.79	4.58	2.40	95.4 (75)
	7	BocTaeg	0.09	0.18	8.55	1.61	0.20	>99.9 (93)
	8	BocTaeg	0.08	0.17	6.53	0.80	0.45	99.0 (91)
į	9	BocTaeg	0.07	0.16	9.26	3.66	0.61	94.8 (86)
35	10	BocTaeg	0.07	0.15	5.32	1.48	0.60	98.8 (93)

(b) Cleavag, Purificati n, and Id ntification of H-[Ta g]₅-Caeg-[Taeg]₄-Lys-NH₇

A portion of protected Boc-[Taeg]₅-Caeg-[Taeg]₄-Lys(Cl2)-BHA resin was treated as described in Example I 5 section (c) to yield about 14.4 mg of crude material upon HF cleavage of 66.9 mg dry H-[Taeg]₅-Caeg-[Taeg]₄-Lys(Cl2)-BHA resin. The main peak at 14.5 min accounted for >50% of the total absorbance. A 100.0 mg portion of the crude product was purified (8 batches; each dissolved in 1 ml H₂O) to give approximately 9.1 mg of 96% pure H-[Taeg]₅-Caeg-[Taeg]₄-Lys-NH₂ (Figure 13b). For (M+H) the calculated m/z value = 2793,8 and the measured m/z value = 2790,6.

EXAMPLE 21

15 Binding of Acr - (Taeg) 10-Lys-MH, to dA10 (Figure 11a)

Acr - (Taeg), -Lys (100 ng) was incubated for 15 min at temperature with 50 cps 5'-[32P]-end-labelled room oligonucleotide [d(GATCCA₁₀G)] in 20 μ l TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 7.4). The sample was cooled in ice (15 20 min) and analyzed by gel electrophoresis in polyacrylamide (PAGE). To 10 μ l of the sample was added 2 μ l 50% glycerol, 5 TBE (TBE = 90 mM Tris-borate, 1 mM EDTA, pH 8.3), and the analysed by PAGE (15% acrylamide, sample was bisacrylamide) in TBE buffer at 4°C. A 10 µl portion of the 25 sample was lyophilized and redissolved in 10 µl 80% formamide, 1 TBE, heated to 90°C (5 min), and analyzed by urea/PAGE (15% acrylamide, 0.5% bisacrylamide, 7 M urea) in TBE. containing DNA bands were visualized by autoradiography using intensifying screens and Agfa Curix RPI X-ray films expos d 30 at -80°C for 2 h.

Oligonucleotides were synthesized on a Biosearch 7500 DNA synthesizer, labelled with $\gamma[^{32}P]$ -ATP (Amersham, 5000 Ci/mmol) and polynucleotide kinase, and purified by PAGE using standard techniques (Maniatis et al. (1986): A laboratory manual, Cold Spring Harbor Laboratories).

EXAMPLE 22

P rmati n f strand displacement complex

A $dA_{10}-dT_{10}$ target sequence contain d within a plasmid DNA sequence was constructed by cloning of two oligonu-5 cleotides $(d(GATCCA_{10}G) + d(GATCCT_{10}G))$ into the BamHI restriction enzyme site of pUC19 using the Eschericia coli JM101 strain by standard techniques (Maniatis et al., 1986). The desired plasmid (designated pT10) was isolated from one of the resulting clones and purified by the alkalin 10 extraction procedure and CsCl centrifugation (Maniatis et al., A 3'-[32P]-end-labelled DNA fragment of 248 bp containing the dA_{10}/dT_{10} target sequence was obtained by cleaving the pT10 DNA with restriction enzymes EcoRI and PvuII, labelling of the cleaved DNA with $\alpha[^{32}P]$ -dATP (4000 15 Ci/mmol, Amersham) using the Klenow fragment of E. coli DNA polymerase (Boehringer Mannheim), and purifying the 248 bp DNA fragment by PAGE (5% acrylamide, 0.06% bisacrylamide, TBE This DNA fragment was obtained with [32P]-endlabelling at the 5'-end by treating the EcoRI-cleaved pT10 20 plasmid with bacterial alkaline phosphatase (Boehring r Mannheim), purifying the plasmid DNA by gel electrophor sis in low melting agarose, and labelling with γ [32 P] ATP and polynucleotide kinase. Following treatment with PvuII, the 248 bp DNA fragment was purified as above.

The complex between $Acr^1-(Taeg)_{10}-Lys-NH_2$ and the 248 bp DNA fragment was formed by incubating 50 ng of $Acr^1-(Taeg)_{10}-Lys-NH_2$ with 500 cps 32 P-labelled 248 bp fragment and 0.5 μ g calf thymus DNA in 100 μ l buffer for 60 min at 37°C.

30 EXAMPLE 23

Probing of strand displacement complex with:

(a) Staphylococcus nuclease (Figure 12b)

The strand displacement complex was formed in 25 mM Tris-HCl, 1 mM MgCl₂, 0.1 mM CaCl₂, pH 7.4 as described above.

35 The comples was treated with Staphylococcus nuclease (B hringer Mannh im) at 750 U/ml f r 5 min at 20°C and the reaction was stopped by addition f EDTA to 25 mM. The DNA

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was precipitated with 2 vols. of ethanol, 2% potassium acetate redissolved in 80% formamide, TBE, heat d to 90°C (5 min), and analyzed by high res lution PAGE (10% acrylamide, 0.3% bisacrylamide, 7 M urea) and autoradiography.

(b) Affinity photocleavage (Figure 12a + 12b)

The complex was formed in TE buffer. A sample contained in an Eppendorf tube was irradiated from above at 300 nm (Philips TL 20 W/12 fluorescent light tube, 24 Jm⁻²s⁻¹) for 30 min. The DNA was precipitated as above, taken up in 1 M piperidine, and heated to 90°C for 20 min. Following lyophilization, the DNA was analysed by PAGE as above.

(c) Potassium permanganate (Figure 12b)

The complex was formed in 100 μ l TE and 5 μ l 20 mM KMnO, was added. After 15 s at 20°C, the reaction was stopped 15 by addition of 50 μ l 1.5 M sodium acetate, pH 7.0, 1 μ M.2-mercaptoethanol. The DNA was precipitated, treated with piperidine and analyzed, as above.

(d) Photofootprinting (Figure 12b)

The complex was formed in 100 µl TE and diazo-linked 20 acridine (0.1 µg/µl) (DHA, Nielsen et al. (1988) Nucl. Acids Res. 16, 3877-88) was added. The sample was irradiated at 365 nm (Philips TL 20 W/09N, 22 Jm⁻²s⁻¹) for 30 min and treated as described for "affinity photocleavage".

(e) S,-nuclease (Pigure 12c)

The complex was formed in 50 mM sodium acetate, 200 mM NaCl, 0.5% glycerol, 1 mM ZnCl₂, pH 4.5 and treated with nuclease S₁ (Boehringer Mannheim) at 0.5 U/ml for 5 min at 20°C. The reaction was stopped and treated further as described under "Staphylococcus nuclease".

30

EXAMPLE 24

N-Benzyloxycarbonyl-N-' (bocaminoethyl) glycine.

Aminoethyl glycine (52.86 g; 0.447 mol) was dissolved in water (900 ml) and dioxane (900 ml) was added. The pH was adjusted to 11.2 with 2N NaOH. While the pH was kept at 11.2, trt-butyl-p-nitrophenyl carbonate (128.4 g; 0.537 mol) was dissolved in dioxane (720 ml) and added dropwise over the

f 2 hours. The pH was kept at 11.2 for at 1 ast three cours mor hours and then I ft with stirring overnight. The y llow solution was cooled to 0°C and the pH was adjusted to 3.5 with 2 N HCl. The mixture was washed with chloroform (4x100 ml). 5 and the pH of the aqueous phase was readjusted to 9.5 with 2 N NaOH at 0°C. Benzyloxycarbonyl chloride (73.5 ml; 0.515 mol) was added over half an hour, while the pH was kept at 9.5 with 2 N NaOH. The pH was adjusted frequently over the next 4 hours, and the solution was left with stirring overnight. 10 On the following day the solution was washed with ether (3x600 ml) and the pH of the solution was afterwards adjusted to 1.5 with 2 N HCl at 0°C. The title compound was isolated by extraction with ethyl acetate (5x1000 ml). The ethyl acetate solution was dried over magnesium sulfate and evaporated to 15 dryness, in vacuo. This afforded 138 g, which was dissolved in ether (300 ml) and precipitated by the addition of petroleum ether (1800 ml). Yield 124.7 g (79%). M.p. 64.5-85 °C. Anal. for $C_{17}H_{22}N_{1}O_{4}$ found(calc.) C: 58.40(57.94); H: 7.02(6.86); N: 7.94(7.95). 'H-NMR (250 MHz, CDCl₃) 7.33 & 7.32 20 (5H, Ph); 5.15 & 5.12 (2H, PhCH); 4.03 & 4.01 (2H, NCH,CO,H); 3.46 (b, 2H, BocNHCH₂CH₂); 3.28 (b, 2H, BocNHCH₂CH₂); 1.43 & 1.40 (9H, 'Bu). HPLC (260 nm) 20.71 min. (80.2%) and 21.57 min. (19.8%). The UV-spectra (200 nm - 300 nm) are identical, indicating that the minor peak consists of Bis-Z-AEG.

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EXAMPLE 25

N'-Boc-aminoethyl glycine ethyl ester.

N-Benzyloxycarbonyl-N'-(bocaminoethyl)glycine (60.0 g; 0.170 mol) and N,N-dimethyl-4-aminopyridine (6.00 g) wer dissolved in absolute ethanol (500 ml), and cooled to 0°C before the addition of DCC (42.2 g; 0.204 mol). The ice bath was removed after 5 minutes and stirring was continued for 2 more hours. The precipitated DCU (32.5 g dried) was removed by filtration and washed with ether (3x100 ml). The combined filtrat was washed successively with diluted potassium hydrogen sulfat (2x400 ml), diluted sodium hydr gencarbonate (2x400 ml) and saturated sodium chl rid (1x400 ml). The

organic phase was filtered, then dried over magnesium sulfate, and evaporated to dryness, in vacuo, which yielded 66.1 g of an oily substance which contained some DCU.

The oil was dissolved in absolute ethanol (600 ml) and 5 was added 10% palladium on carbon (6.6 g) was added. solution was hydrogenated at atmospheric pressure, where the reservoir was filled with 2 N sodium hydroxide. hours, 3.3 L was consumed out of the theoretical 4.2 L. The reaction mixture was filtered through celite and evaporated 10 to dryness, in vacuo, affording 39.5 g (94%) of an oily substance. A 13 g portion of the oily substance was purified by silica gel (600 g SiO₂) chromatography. After elution with 300 ml 20% petroleum ether in methylene chloride, the title compound was eluted with 1700 ml of 5% methanol in methylene The solvent was removed from the fractions with 15 chloride. satisfactory purity, in vacuo and the yield was 8.49 g. Alternatively 10 q of the crude material was purified by Kugel Rohr distillation. H-NMR (250 MHz, CD,OD); 4.77 (b. s, NH); 4.18 (q, 2H, MeCH-); 3.38 (s, 2H, NCH-CO-Et); 3.16 (t, 2H, 20 BocNHCH,CH,); 2.68 (t, 2H, BocNHCH,CH.); 1.43 (s, 9H, 'Bu) and 1.26 (t, 3H, CH₁) "C-NMR 171.4 (<u>C</u>OEt); 156.6 (CO); 78.3 $((CH_1),C)$; 59.9 (CH_2) ; 49.0 (CH_2) ; 48.1 (CH_2) ; 39.0 (CH_2) ; 26.9 (CH₂) and 12.6 (CH₃).

25 EXAMPLE 26

N'-Boc-aminoethyl glycine methyl ester.

The above procedure was used, with methanol being substituted for ethanol. The final product was purified by column purification.

30

EXAMPLE 27

1-(Boc-aeg) thymine ethyl ester.

N'-Boc-aminoethyl glycine ethyl ester (13.5 g; 54.8 mmol), DhbtOH (9.84 g; 60.3 mmol) and 1-carboxymethyl thymine (11.1 g; 60.3 mmol) were diss lved in DMF (210 ml). Methylene chloride (210 ml) then was added. The solution was cooled t 0°C in an ethanol/ice bath and DCC (13.6 g; 65.8

mmol) was add d. The ice bath was rem ved aft r 1 h ur and stirring was continued f r another 2 h urs at ambient temperature. The precipitated DCU was removed by filtration and washed twice with methylene chloride (2 x 75 ml). To the 5 combined filtrate was added more methylene chloride (650 ml). The solution was washed successively with diluted.sodium hydrogen carbonate (3 x 500 ml), diluted potassium hydrog n sulfate (2 x 500 ml), and saturated sodium chloride (1 x 500 ml). Some precipitate was removed from the organic phase by 10 filtration, The organic phase was dried over magnesium sulfate and evaporated to dryness, in vacuo. The oily residue was dissolved in methylene chloride (150 ml), filtered, and the title compound was precipitated by the addition petroleum ether (350 ml) at 0°C. The 15 chloride/petroleum ether procedure was repeated once. This afforded 16.0 g (71%) of a material which was more than 99% pure by HPLC.

EXAMPLE 28

20 1-(Boc-aeg)thymine.

The material from above was suspended in THF (194 ml, gives a 0.2 M solution), and 1 M aqueous lithium hydroxide (116 ml) was added. The mixture was stirred for 45 minutes at ambient temperature and then filtered to remove residual 25 DCU. Water (40 ml) was added to the solution which was then washed with methylene chloride (300 ml). Additional water (30 ml) was added, and the alkaline solution was washed once more with methylene chloride (150 ml). The aqueous solution was cooled to 0°C and the pH was adjusted to 2 by the dropwis 30 addition of 1 N HCl (approx. 110 ml). The title compound was extracted with ethyl acetate (9 x 200 ml), the combin d extracts were dried over magnesium sulfate and were evaporated to dryness, in vacuo. The residue was evaporated once from methanol, which after drying overnight afforded a colorless 35 glassy solid. Yield 9.57 g (64 %). HPLC > 98% R_{τ} =14.8 min. Anal. for $C_{16}H_{24}N_4O_7^{\bullet}0.25 H_7O$ F und (calc.) C: 49.29(49.42); H: 6.52(6.35); N: 14.11(14.41). Due t the limited rotati n around the secondary amide, several f the signals were doubled in the ratio 2:1 (indicated in the list by mj. for major and mi. for minor). H-NMR (250 MHz, DMSO-d₆): 12.75 (b.s., 1H, CO₂H); 11.28 (s, "1H", mj., imide NH); 11.26 (s, 5 "1H", mi., imide NH); 7.30 (s, "1H", mj., T H-6); 7.26 (s, "1H", mi., T H-6); 6.92 (b.t., "1H", mj., BocNH); 6.73. (b.t., "1H", mi., BocNH); 4.64 (s, "2H", mj., CH₂CON); 4.46 (s, "2H", mj., CH₂CON); 4.46 (s, "2H", mj., CH₂CO₂H); 3.97 (s, "2H", mj., CH₂CO₂H); 3.63-3.01 (unresolved m, includes water, 10 CH₂CH₂); 1.75 (s, 3H, CH₃) and 1.38 (s, 9H, ⁵Bu).

EXAMPLE 29

N'-Benzyloxycarbonyl-1-(Boc-aeg) cytosine.

N'-Boc-aminoethyl glycine ethyl ester (5.00 g; 20.3 15 mmol), DhbtOH (3.64 g; 22.3 mmol) and N-benzyloxycarbonyl-1carboxymethyl cytosine (6.77 g; 22.3 mmol) were suspended in DMF (100 ml). Methylene chloride (100 ml) then was added. The solution was cooled to 0°C and DCC (5.03 g; 24.4 mmol) was added. The ice bath was removed after 2 h and stirring was 20 continued for another hour at ambient temperature. reaction mixture then was evaporated to dryness, in vacuo. The residue was suspended in ether (100 ml) and stirred vigorously for 30 min. The solid material was isolated by filtration and the ether wash procedure was repeated twice. 25 The material was then stirred vigorously for 15 min with dilute sodium hydrogencarbonate (aprox. 4% solution, 100 ml), filtered and washed with water. This procedure was then repeated once, which after drying left 17.0 g of yellowish solid material. The solid was then boiled with dioxane (200 30 ml) and filtered while hot. After cooling, water (200 ml) was - added. The precipitated material was isolated by filtration, washed with water, and dried. According to HPLC (observing at 260 nm) this material has a purity higher than 99%, besides the DCU. The ester was then suspended in THF (100 ml), cooled 35 to 0°C, and 1 N LiOH (61 ml) was added. After stirring for 15 minutes, the mixture was filter d and the filtrate was washed with methylene chloride (2 x 150 ml). The alkaline WO 92/20702 PCT/EP92/01219

solution then was cooled t 0°C and the pH was adjusted to 2.0 with 1 N HCl. The title compound was isolated by filtrati n and was washed nce with water, leaving 11.3 g of a white powder after drying. The material was suspended in methylen 5 chloride (300 ml) and petroleum ether (300 ml) was added. Filtration and wash afforded 7.1 g (69%) after drying, HPLC showed a purity of 99% R = 19.5 min, and a minor impurity at 12.6 min (approx. 1%) most likely the Z-de protected monomer. Anal. for $C_2H_2N_5O_4$ found(calc.) C: 54.16(54.87); H: 5.76(5.81) 10 and N: 13.65(13.91). H-NMR (250 MHz, DMSO-d₄). 10.78 (b.s. 1H, COH); 7.88 (2 overlapping dublets, 1H, Cyt H-5); 7.41-7.32 (m, 5H, Ph); 7.01 (2 overlapping doublets, 1H, Cyt H-6); 6.94 & 6.78 (unres. triplets, 1H, BocNH); 5.19 (s, 2H, PhCH); 4.81 & 4.62 (s, 2H, CH.CON); 4.17 & 3.98 (s, 2H, CH.CO.H); 15 3.42-3.03 (m, includes water, CH.CH.) and 1.38 & 1.37 (s, 9H, ¹⁰C-NMR. 150.88; 128.52; 128.18; 127.96; 93.90; 66.53; IR: Frequency in cm⁻¹ (intensity). 3423 49.58 and 28.22. (26.4), 3035 (53.2), 2978(41.4), 1736(17.3), 1658(3.8), 1563(23.0), 1501(6.8) and 1456 (26.4).

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EXAMPLE 30

9-Carboxymethyl adenine ethyl ester.

Adenine (10.0 g, 74 mmol) and potassium carbonate (10.29 g, 74.0 mmol) were suspended in DMF and ethyl 25 bromoacetate (8.24 ml, 74 mmol) was added. The suspension was stirred for 2.5 h under nitrogen at room temperature and then filtered. The solid residue was washed three times with DMF (10 ml). The combined filtrate was evaporated to dryness, in vacuo. The yellow-orange solid material was poured into water 30 (200 ml) and 4 N HCl was added to pH≈6. After stirring at 0°C for 10 min, the solid was filtered off, washed with water, and recrystallized from 96% ethanol (150 ml). The title comp und was isolated by filtration and washed thoroughly with ether. Yield 3.4 g (20%). M.p. 215.5-220°C. Anal. for CoH, NoO, 35 found(calc.): C: 48.86(48.65); H: 5.01(4.91); N: 31.66(31.42). $^{1}H-NMR$ (250 MHz; DMSO- d_{A}): (s, 2H, H-2 & H-8), 7.25 (b. s., 2H, NH_2), 5.06 (s, 2H, NCH_2), 4.17 (q, 2H, J=7.11 Hz, OCH_2) and 1.21 (t, 3H, J=7.13 Hz, NCH₂). ¹³C-NMR. 152.70, 141.30, 61.41, 43.97 and 14.07. FAB-MS. 222 (MH+). IR: Frequency in cm⁻¹ (intensity). 3855 (54.3), 3274(10.4), 3246(14.0), 3117(5.3), 2989(22.3), 2940(33.9), 2876(43.4), 2753(49.0), 2346(56.1), 2106(57.1), 1899(55.7), 1762(14.2), 1742(14.2), 1742(1.0), 1671(1.8), 1644(10.9), 1606(0.6), 1582(7.1), 1522(43.8), 1477(7.2), 1445(35.8) and 1422(8.6). The position f alkylation was verified by X-ray crystallography on crystals, which were obtained by recrystallization from 96% ethanol.

Alternatively, 9-carboxymethyl adenine ethyl ester can be prepared by the following procedure. To a suspension f adenine (50.0 g, 0.37 mol) in DMF (1100 ml) in 2 L thre-necked flask equipped with a nitrogen inlet, a mechanical stirrer and a dropping funnel was added 16.4 g (0.407 mol) haxane washed sodium hydride- mineral oil dispersion. The mixture was stirred vigorously for 2 hours, whereafter ethy bromacetate 75 ml, 0.67 mol) was added dropwise over the course of 3 hours. The mixture was stirred for one additional hour, whereafter tlc indicated complete conversion of adenine.

The mixture was evaporated to dryness at 1 mmHg and water (500 ml) was added to the oily residue which caused crystallisation

of the title compound. the solid was recrystallised from 06% ethanol (600 ml). Yield after drying 53.7 (65.6%). HPLC (215

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EXAMPLE 31

nm) purity > 99.5%.

M'Benzyloxycarbonyl-9-carboxymethyl adenine ethyl ester.

9-Carboxymethyladenine ethyl ester (3.40 g, 15.4 mmol) was dissolved in dry DMF (50 ml) by gentle heating, cooled to added to solution ٥f N-ethyl-30 20°C. and zyloxycarbonylimidazole tetrafluoroborate (62 mmol) methylene chloride (50 ml) over a period of 15 min with icecooling. Some precipitation was observed. The ice bath was removed and the solution was stirred overnight. The reaction 35 mixture was treated with saturated sodium hydrogen carbonate Aft r stirring for 10 min, the phases were sepafated and the organic phase was washed successively with

volume of water, dilute potassium hydrogen sulfat (twice), and with saturat d sodium chloride. The s luti n was dried over magnesium sulfate and evaporated to dryness, in vacuo, which afforded 11 g of an oily material. The material 5 was dissolved in methylene chloride (25 ml), cooled to 0°C, and precipitated with petroleumeum ether (50 ml). This procedure was repeated once to give 3.45 g (63%) of the title compound. M.p. 132-35°C. Analysis for C17H17N5O4 found (calc.): C: 56.95(57.46); H: 4.71(4.82); N: 19.35(19.71). 1H-10 NMR (250 MHz; CDCl3): 8.77 (s, 1H, H-2 or H-8); 7.99 (s, 1H, H-2 or H-8); 7.45-7.26 (m, 5H, Ph); 5.31 (s, 2H, N-CH₂); 4.96 (s, 2H, Ph-C H_2); 4.27 (q, 2H, J=7.15 Hz, C H_3 C H_4) and 1.30 (t, 3H, J=7.15 Hz, CH, CH_3). "C-NMR: 153.09; 143.11; 128.66; 67.84; 62.51; 44.24 and 14.09. FAB-MS: 356 (MH+) and 312 15 (MH+-CO₂). IR: frequency in cm⁻¹ (intensity). 3423 (52.1); 3182 (52.8); 3115(52.1); 3031(47.9); 2981(38.6); 1747(1.1); 1617(4.8); 15.87(8.4); 1552(25.2); 1511(45.2); 1492(37.9); 1465(14.0) and 1413(37.3).

20 EXAMPLE 32

N'Benzyloxycarbonyl-9-carboxymethyl adenine.

N°-Benzyloxycarbonyl-9-carboxymethyladenine ethyl ester (3.20 g; 9.01 mmol) was mixed with methanol (50 ml) cooled to 0°C. Sodium Hydroxide Solution (50 ml; 2N) was added, whereby 25 the material quickly dissolved. After 30 min at 0°C, th alkaline solution was washed with methylene chloride (2x50ml). The aqueous solution was brought to pH 1.0 with 4 N HCl at 0°C, whereby the title compound precipitated. The yield after filtration, washing with water, and drying was 3.08 g (104%). 30 The product contained salt and elemental analysis reflect d that. Anal. for $C_{15}H_{13}N_5O_4$ found(calc.): C: 46.32(55.05); H: 4.24(4.00); N: 18.10(21.40) and C/N: 2.57(2.56). H-NMR(250 MH2; $DMSO-d_6$): 8.70 (s, 2H, H-2 and H-8); 7.50-7.35 (m, 5H, Ph); 5.27 (s, 2H, N-CH₂); and 5.15 (s, 2H, Ph-CH₂). 13 C-NMR. 35 168.77, 152.54, 151.36, 148.75, 145.13, 128.51, 128.17,127.98, 66.76 and 44.67.IR (KBr) 3484(18.3); 3109(15.9); 3087(15.0); 2966(17.1); 2927(19.9); 2383(53.8); 1960(62.7); 1739(2.5);

1688(5.2); 1655(0.9); 1594(11.7); 1560(12.3); 1530(26.3); 1499(30.5); 1475(10.4); 1455(14.0); 1429(24.5) and 1411(23.6). FAB-MS: 328 (MH+) and 284 (MH+-CO₂). HPLC (215 nm, 260 nm) in system 1: 15.18 min, minor impurities all less than 2%.

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EXAMPLE 33

M'-Benzyloxycarbonyl-1-(Boc-aeg)adenine ethyl ester.

N'-Boc-aminoethyl glycine ethyl ester (2.00 g; 8.12 mmol), DhbtOH (1.46 g; 8.93 mmol) and N'-benzyloxycarbonyl-9-10 carboxymethyl adenine (2.92 g; 8.93 mmol) were dissolved in DMF (15 ml). Methylene chloride (15 ml) then was added. The solution was cooled to 0°C in an ethanol/ice bath. DCC (2.01 g; 9.74 mmol) was added. The ice bath was removed after 2.5 h and stirring was continued for another 1.5 hour at ambient 15 temperature. The precipitated DCU was removed by filtration and washed once with DMF (15 ml), and twice with methylene chloride (2 x 15 ml). To the combined filtrate was added more methylene chloride (100 ml). The solution was washed. successively with dilute sodium hydrogen carbonate (2 x 100 20 ml), dilute potassium hydrogen sulfate (2 x 100 ml), and saturated sodium chloride (1 x 100 ml). The organic phase was evaporated to dryness, in vacuo, which afforded 3.28 g (73%) of a yellowish oily substance. HPLC of the raw product show d a purity of only 66% with several impurities, both more and 25 less polar than the main peak. The oil was dissolved in absolute ethanol (50 ml) and activated carbon was added. After stirring for 5 minutes, the solution was filtered. The filtrate was mixed with water (30 ml) and was left with stirring overnight. The next day, the white precipitate was 30 removed by filtration, washed with water, and dried, affording _ 1.16 g (26%) of a material with a purity higher than 98% by HPLC. Addition of water to the mother liquor afforded another 0.53 g with a purity of approx. 95%. Anal. for C2H2N7O7°H2O found(calc.) C: 55.01(54.44; H: 6.85(6.15) 35 16.47(17.09). 'H-NMR (250 MHz, CDCl₃) 8.74 (s, 1H, Ade H-2); 8.18 (b. s, 1H, ZNH); 8.10 & 8.04 (s, 1H, H-8); 7.46-7.34 (m, 5H, Ph); 5.63 (unr s. t, 1H, BocNH); 5.30 (s, 2H, PhCH2); 5.16

£ 5.00 (s, 2H, CH_CON); 4.29 £ 4.06 (s, 2H, CH_CO,H); 4.20 (q, 2H, OCH_CH_); 3.67-3.29 (m, 4H, CH_CH_); 1.42 (s, 9H, 'Bu) and 1.27 (t, 3H, OCH_CH_). The spectrum shows traces of ethanol and DCU.

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EXAMPLE 34

N'-Benzyloxycarbonyl-1-(Boc-aeg) adenine.

N'-Benzyloxycarbonyl-1-(Boc-aeg) adenine ethyl ester (1.48 g; 2.66 mmol) was suspended in THF (13 ml) and the 10 mixture was cooled to 0°C. Lithium hydroxide (8 ml; 1 N) was added. After 15 min of stirring, the reaction mixture was filtered, extra water (25 ml) was added, and the solution was washed with methylene chloride (2 x 25 ml). The pH of th aqueous solution was adjusted to pH 2.0 with 1 N HCl. The 15 precipitate was isolated by filtration, washed with water, and dried, and drief affording 0.82 g (58%). The product reprecipitated twice with methylene chloride/petroleum ether, 0.77 g (55%) after drying. M.p. 119°C (decomp.) Anal. for $C_{24}H_{24}N_{1}O_{1}^{\circ}H_{2}O$ found(calc.) C: 53.32(52.84); H: 5.71(5.73); N: 20 17.68(17.97). FAB-MS. 528.5 (MH+). 'H-NMR (250 MHz, DMSO-d4). 12.75 (very b, 1H, CO₂H); 10.65 (b. s, 1H, ZNH); 8.59 (d, 1H, J=2.14 Hz, Ade H-2); 8.31 (s, 1H, Ade H-8); 7.49-7.31 (m, 5H, Ph); 7.03 & 6.75 (unresol. t, 1H, BocNH); 5.33 & 5.16 (s, 2H, CH,CON); 5.22 (s, 2H, PhCH); 4.34-3.99 (s, 2H, CH,CO,H); 3.54-25 3.03 (m's, includes water, CH.CH.) and 1.39 & 1.37 (s, 9H, "C-NMR. 170.4; 166.6; 152.3; 151.5; 149.5; 145.2; 128.5; 128.0; 127.9; 66.32; 47.63; 47.03; 43.87 and 28.24.

EXAMPLE 35

30 2-Amino-6-chloro-9-carboxymethylpurine.

To a suspension of 2-amino-6-chloropurine (5.02 g; 29.6 mmol) and potassium carbonate (12.91 g; 93.5 mmol) in DMF (50 ml) was added bromoacetic acid (4.70 g; 22.8 mmol). The mixture was stirred vigorously for 20 h. under nitrogen.

35 Wat r (150 ml) was add d and th s luti n was filtered through Celite to give a cl ar yellow solution. The solution was acidifi d t a pH f 3 with 4 N hydr chl ric acid. The

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precipitate was filtered and dried, in vacuo, ver sicapent. Yield (3.02 g; 44.8%). 'H-NMR(DMSO-d6): d = 4.88 ppm (s,2H); 6.95 (s,2H); 8.10 (s,1H).

5 EXAMPLE 36

2-Amino-6-benzyloxy-9-carboxymethylpurine.

Sodium (2.0 g; 87.0 mmol) was dissolved in benzyl alcohol (20 ml) and heated to 130°C for 2 h. After cooling to 0°C, a solution of 2-amino-6-chloro-9-carboxymethylpurine 10 (4.05 g; 18.0 mmol) in DMF (85 ml) was slowly added, and the resulting suspension stirred overnight at 20°C. hydroxide solution (1N, 100 ml) was added and the clear solution was washed with ethyl acetate (3 x 100 ml). water phase then was acidified to a pH of 3 with 4 N 15 hydrochloric acid. The precipitate was taken up in ethyl acetate (200 ml), and the water phase was extracted with ethyl acetate (2 x 100 ml). The combined organic phases were washed with saturated sodium chloride solution (2 x 75 ml), dried with anhydrous sodium sulfate, and taken to dryness by 20 evaporation, in vacuo. The residue was recrystallized from ethanol (300 ml). Yield after drying, in vacou, sicapent: 2.76 g (52%). M.p. 159-65°C. Anal. (calc., found) C(56.18; 55.97), H(4.38; 4.32), N(23.4; 23.10). H-NMR (DMSOd₂): 4.82 ppm.'(s,2H); 5.51 (s,2H); 6.45 (s,2H); 7.45 (m,5H); 25 7.82 (s,1H).

EXAMPLE 37

N-([2-Amino-6-bensyloxy-purine-9-yl]-acetyl)-N-(2-Boc-aminoethyl)-glycine [BocGaeg-OH monomer].

2-Amino-6-benzyloxy-9-carboxymethyl-purine (0.50 g;
1.67 mmol), methyl-N(2-[tert-butoxycarbonylamino]ethyl)glycinate (0.65 g; 2.80 mmol), diisopropylethyl amine (0.54 g; 4.19 mmol), and bromo-tris-pyrrolidino-phosphoniumhexafluoro-phosphate (PyBroP®) (0.798 g; 1.71 mmol) were

35 stirred in DMF (2 ml) f r 4 h. The clear solution was poured int an ice-co led solution of s dium hydrogen carbonate (1 N; 40 ml) and extracted with ethyl acetate (3 X 40 ml). The

organic layer was washed with potassium hydrogen sulfate solution (1 N; 2 X 40 ml), s dium hydr gen carbonate (1 N; 1 X 40 ml) and saturated sodium chloride solution (60 ml). After drying with anhydrous sodium sulfate and evaporation, 5 in vacuo, the solid residue was recrystallized from ethyl acetate/hexane (20 ml (2:1)) to give the methyl ester in 63% yield (MS-FAB 514 (M+1). Hydrolysis was accomplished by dissolving the ester in ethanol/water (30 ml (1:2)) containing conc. sodium hydroxide (1 ml). After stirring for 2 h, the 10 solution was filtered and acidified to a pH of 3, by th addition of 4 N hydrochloric acid. The title compound was Yield: 370 mg (72% for the obtained by filtration. hydrolysis). Purity by HPLC was more than 99%. Due to th limited rotation around the secondary amide several of th 15 signals were doubled in the ratio 2:1 (indicated in the list by mj. for major and mi. for minor). 1H-NMR(250, MHz, DMSO d_6): d = 1.4 ppm. (s,9H); 3.2 (m,2H); 3.6 (m,2H); 4.1 (s, mj.,CONRCH2COOH); 4.4 (s, mi., CONRCH2COOH); 5.0 (s, mi., Gua- $CH_2CO-)$; 5.2 (s, mj., $Gua-CH_2CO)$; 5.6 (s,2H); 6.5 (s,2H); 6.9 20 (m, mi., BocNH); 7.1 (m, mj., BocNH); 7.5 (m.,3H); 7.8 (s,1H); 12,8 (s;1H). "C-NMR. 170.95; 170.52; 167.29; 166.85; 160.03; 159.78; 155.84; 154.87; 140.63; 136.76; 128.49; 128.10; 113.04; 78.19; 77.86; 66.95; 49.22; 47.70; 46.94; 45.96; 43.62; 43.31 and 28.25.

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EXAMPLE 38

3-Boc-amino-1,2-propanediol.

3-Amino-1,2-propanediol (40.00 g, 0.440 mol, 1.0 q.)
was dissolved in water (1000 ml) and cooled to 0°C. Di-t rt30 butyl dicarbonate (115.0 g, 0.526 mol, 1.2 eq.) was add d in
one portion. The reaction mixture was heated to room
temperature on a water bath during stirring. The pH was
maintained at 10.5 with a solution of sodium hydroxide (17.56
g, 0.440 mol, 1.0 eq.) in water (120 ml). When the addition
35 of aqueous sodium hydroxide was completed, the reaction
mixtur was stirr d ov rnight at ro m t mperature.
Subsequ ntly, thyl acetate (750 ml) was added to the reaction

mixture, f llowed by cooling to 0°C. The pH was adjusted to 2.5 with 4 N sulphuric acid with vigor us stirring. phases were separat d and the water phas was washed with additional ethyl acetate (6x350 ml). The volume of the 5 organic phase was reduced to 900 ml by evaporation under reduced pressure. The organic phase then was washed with a saturated aqueous solution of potassium hydrogen sulfat diluted to twice its volume (1x1000 ml) and with saturated aqueous sodium chloride (1x500 ml). The organic phase was 10 dried (MgSO,) and evaporated under reduced pressure to yield 50.12 g (60%) of the title compound. The product could be solidified by evaporation from methylene chloride and subsequent freezing. 'H-NMR (CDCl,/TMS): d = 1.43 (s, 9H, $Me_{1}C$), 3.25 (m, 2H, CH_{2}), 3.57 (m, 2H, CH_{2}), 3.73 (m, 1H, CH). 15 "C-NMR (CDCl₁/TMS): d = 28.2 (Me₃C), 42.6 (CH₂), 63.5, 71.1 (CH₂OH, CHOH), 79.5 (Me₃C), 157.0 (C=0).

EXAMPLE 39

2-(Boc-amino)ethyl-L-alanine methyl ester.

3-Boc-amino-1,2-propanediol (20.76 g, 0.109 mol, 1 eq.) was suspended in water (150 ml). Potassium m-periodate (24.97 g, 0.109 mol, 1 eq.) was added and the reaction mixture was stirred for 2 h at room temperature under nitrogen. The reaction mixture was filtered and the water phase extracted with chloroform (6x250 ml) The organic phase was dried (MgSO₂) and evaporated to afford an almost quantitative yield of Bocaminoacetaldehyde as a colourless oil, which was used without further purification in the following procedure.

Palladium-on-carbon (10%, 0.8 g) was added to MeOH (250 ml) under nitrogen with cooling (0°C) and vigorous stirring. Anhydrous sodium acetate (4.49 g, 54.7 mmol, 2 eqv) and Lalanine methyl ester, hydrochloride (3.82 g, 27.4 mmol, 1 eqv) were added. Boc-aminoacetaldehyde (4.79 g, 30.1 mmol, 1.1 eqv) was dissolved in MeOH (150 ml) and added to the reaction mixture. The reaction mixture was hydrogenated at atmospheric pr ssure and r om temperature until hydrogen uptake had ceas d. The reaction mixture was filtered through celite,

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which was washed with additional MeOH. The MeOH was removed under reduced pr ssur. The r sidue was suspended in water (150 ml) and pH adjusted to 8.0 by dropwise addition of 0.5 N NaOH with vigorous stirring. The water phase was extract d 5 with methylene chloride (4x250 ml). The organic phase was dried (MgSO₄), filtered through celite, and evaporated under reduced pressure to yield 6.36 g (94%) of the title compound as a clear, slightly yellow oil. MS (FAB-MS): m/z (%) = 247 (100, M+1, 191 (90), 147 (18). H-NMR (250 MHz, CDCl₃). 1.18 10 (d, J=7.0 Hz, 3H, Me), 1.36 (s, 9H, Me₃C), 1.89 (b, 1H, NH), 2.51 (m, 1H, CH₂), 2.66 (m, 1H, CH₂), 3.10 (m, 2H, CH₂), 3.27 (q, J=7.0 Hz, 1H, CH), 3.64 (s, 3H, OMe), 5.06 (b, 1H, carbamate NH). G-NMR. d = 18.8 (Me), 28.2 (Me₃C), 40.1, 47.0 (CH₂), 51.6 (OMe), 56.0 (CH), 155.8 (carbamate C=O), 175.8 15 (ester C=O).

EXAMPLE 40

N-(Boc-aminosthyl)-N-(1-thyminylacetyl)-L-alanine methyl ester.

To a solution of Boc-aminoethyl-(L)-alanine methyl 20 ester (1.23 g, 5.0 mmol) in DMF (10 ml) was added Dhbt-OH (0,90 g, 5.52 mmol) and 1-thyminylacetic acid (1.01 g, 5.48 When the 1-thyminylacetic acid was dissolved, dichloromethane (10 ml) was added and the solution was cooled 25 on an ice bath. After the reaction mixture had reached 0°C, DCC (1.24 g, 6.01 mmol) was added. Within 5 min after th addition, a precipitate of DCU was seen. After a further 5 min, the ice bath was removed. Two hours later, TLC analysis showed the reaction to be finished. The mixture was filter d 30 and the precipitate washed with dichloromethane (100ml). The resulting solution was extracted twice with 5% sodium hydrogen carbonate (150 ml) and twice with saturated potassium hydr g n sulfate (25 ml) in water (100 ml). After a final extracti n with saturated sodium chloride (150 ml), the solution was 35 dried with magnesium sulfate and evaporated to give a whit The foam was purifi d by c lumn chr matography on silica gel using dichlor m thane with a methanol gradient as eluent. This yielded a pure compound (>99% by HPLC) (1.08 g, 52.4%). FAB-MS: 413 (M+1) and 431 (M+1 + water). H-NMR (CDCl₃): 4.52 (s, 2 H, CH'₂); 3,73 (s, 3 H, OMe); 3.2-3.6 (m, 4 H, ethyl CH₂'s); 1.90 (s, 3 H, Me in T); 1.49 (d, 3 H, M in Ala, J=7.3 Hz); 1.44 (s, 9 H, Boc).

EXAMPLE 41

N-(Boc-aminoethyl)-N-(1-thyminylacetyl)-L-alanine.

The methyl ester of the title compound (2.07 g, 5.02 10 mmol) was dissolved in methanol (100 ml), and cooled on an ice 2 M sodium hydroxide (100 ml) was added. stirring for 10 min, the pH of the mixture was adjusted to 3 with 4 M hydrogen chloride. The solution was subsequently extracted with ethyl acetate (3 x 100 ml). The combined 15 organic extracts were dried over magnesium sulfate. After evaporation, the resulting foam was dissolved in ethyl acetate (400 ml) and a few ml of methanol to dissolve the solid material. Petroleum ether then was added until precipitation started. After standing overnight at -20°C, the precipitate 20 was removed by filtration. This gave 1.01 g (50.5%) of pure compound (>99% by HPLC). The compound can be recrystallized from 2-propanol. FAB-MS: 399 (M+1). H-NMR (DMSO-d₆): 11.35 (s, 1 H, COO); 7.42 (s, 1 H, H₆); 4.69 (s, 2 H, CH₇); 1.83 (s, 3 H, Me in T); 1.50-1.40 (m, 12 H, Me in Ala + Boc).

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EXAMPLE 42

(a) N-(Boc-aminoethyl)--N-(1-thyminylacetyl)-palanine methyl ester.

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To a solution of Boc-aminoethyl alanine methyl ester 30 (2.48 g, 10.1 mmol) in DMF (20 ml) was added Dhbt-OH (1.80 g, 11.0 mmol) and thyminylacetic acid (2.14 g, 11.6 mmol). After dissolution of the 1-thyminylacetic acid, methylene chloride (20 ml) was added and the solution cooled on an ice bath. When the reaction mixture had reached 0°C, DCC (2.88 g, 14.0 mmol) was added. Within 5 min after the addition a pr cipitat f DCU was s en. Aft r 35 min the ice bath was removed. The reaction mixture was filtered 3.5 h later and

the pr cipitate washed with methylene chloride (200 ml). The resulting solution was extracted twice with 5% sodium hydrogen carbonate (200 ml) and twice with saturated potassium hydrogen sulfate in water (100 ml). After a final extraction with saturated sodium chloride (250 ml), the solution was dried with magnesium sulfate and evaporated to give an oil. The oil was purified by short column silica gel chromatography using methylene chloride with a methanol gradient as eluent. This yielded a compound which was 96% pure according to HPLC (1.05 g, 25.3%) after precipitation with petroleum ether. FAB-MS: 413 (M+1). H-NMR (CDCl₃): 5.64 (t, 1 H, BocNH, J=5.89 Hz); 4.56 (d, 2 H, CH'₂); 4.35 (q, 1 H, CH in Ala, J=7.25 Hz); 3.74 (s, 3 H, OMe); 3.64-3.27 (m, 4 H, ethyl H's); 1.90 (s, 3 H, Me in T); 1.52-1.44 (t, 12 H, Boc+Me in Ala).

(b) N-(Boc-aminoethyl)-N-(1-thyminylacetyl)-D-alanin 15 The methyl ester of the title compound (1.57 g, 3.81 mmol) was dissolved in methanol (100 ml) and cooled on an ice Sodium hydroxide (100 ml; 2 M) was added. stirring for 10 min the pH of the mixture was adjusted t 3 20 with 4 M hydrogen chloride. The solution then was extract d The combined organic with ethyl acetate (3 x 100 ml). extracts were dried over magnesium sulfate. After evaporation, the oil was dissolved in ethyl acetate (200 ml). Petroleum ether was added (to a total volume of 600 ml) until 25 precipitation started. After standing overnight at -20°C, th precipitate was removed by filtration. This afforded 1.02 g (67.3%) of the title compound, which was 94% pure according to HPLC. FAB-MS: 399 (M+1). H-NMR: 11.34 (s, 1 H, COOH); 7.42 (s, 1 H, H₆); 4.69 (s, 2 H, CH₂); 4.40 (q, 1 H, CH in 30 Ala, J=7.20 Hz); 1.83 (s, 3 H, Me in T); 1.52-1.40 (m, 12 H, Boc + Me in Ala).

EXAMPLE 43

N-(N'-Boc-3'-aminopropyl)-N-[(1-thyminyl)acetyl]glycine m thyl sester.

N-(N'-Boc-3'-aminopropyl) glycine methyl ester (2.84 g, 0.0115 mol) was dissolved in DMF (35 ml), follow d by addition

of DhbtOH (2.07 g, 0.0127 mol) and 1-thyminylacetic acid (2.34 g, 0.0127 mol). Methylene chl ride (35 ml) was added and the mixture cooled t 0°C on an ice bath. After addition of DCC (2.85 g, 0.0138 mol), the mixture was stirred at 0°C for 2 h, 5 followed by 1 h at room temperature. The precipitated DCU was removed by filtration, washed with methylene chloride (25 ml), and a further amount of methylene chloride (150 ml) was added to the filtrate. The organic phase was extracted with sodium hydrogen carbonate (1 volume saturated diluted with 1 volume 10 water, 6 x 250 ml), potassium sulfate (1 volume saturated diluted with 4 volumes water, 3 x 250 ml), and saturated aqueous sodium chloride (1 x 250 ml), dried over magnesium sulfate, and evaporated to dryness, in vacuo. residue was suspended in methylene chloride (35 ml) and 15 stirred for 1h. The precipitated DCU was removed by filtration and washed with methylene chloride (25 ml). The filtrate was evaporated to dryness, in vacuo, and the residue purified by column chromatography on silica gel, eluting with a mixture of methanol and methylene chloride (gradient from 20 3-7% methanol in methylene chloride). This afforded the title compound as a white solid (3.05 g, 64%). M.p.. 76-79°C (decomp.). Anal. for $C_{18}H_{28}N_4O_7$, found (calc.) C: 52.03 (52.42) H: 6.90 (6.84) N: 13.21 (13.58). The compound showed satisfactory ¹H and ¹³C-NMR spectra.

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EXAMPLE 44

N-(N'-Boc-3'-aminopropyl)-N-[(1-thyminyl)acetyl]glycine.

N-(N'-Boc-3'-aminopropyl)-N-[(1-thyminyl)acetyl]glycine methyl ester (3.02 g, 0.00732 mol) was dissolved in methanol (25 ml) and stirred for 1.5 h with 2 M sodium hydroxide (25 ml). The methanol was removed by evaporation, in vacuo, and pH adjusted to 2 with 4 M hydrochloric acid at 0°C. The product was isolated as white crystals by filtration, washed with water (3 x 10 ml), and dried over sicapent, in vacuo.

35 Yield 2.19 g (75%). Anal. for C₁₇H₂₆N₄O₇, H₂O, found (calc.) C: 49.95 (49.03) H: 6.47 (6.29) N: 13.43 (13.45). The compound showed satisfact ry ¹H and ¹³C-NMR spectra.

3-(1-Thyminy1)-propan ic acid m thyl ester.

Thymine (14.0 g, 0.11 mol) was suspended in methanol. Methyl acrylate (39.6 ml, 0.44 mol) was added, along with 5 catalytic amounts of sodium hydroxide. The solution was refluxed in the dark for 45 h, evaporated to dryness, in vacuo, and the residue dissolved in methanol (8 ml) with heating. After cooling on an ice bath, the product was precipitated by addition of ether (20 ml), isolated by filtration, washed with ether (3 x 15 ml), and dried over sicapent, in vacuo. Yield 11.23 g (48%). M.p. 112-119°C. Anal. for C₉H₁₂N₂O₄, found (calc.) C: 51.14 (50.94) H: 5.78 (5.70) N: 11.52 (13.20). The compound showed satisfactory ¹H and ¹³C-NMR spectra.

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EXAMPLE 46

3-(1-Thyminyl)-propanoic acid.

3-(1-Thyminyl)-propanoic acid methyl ester (1.0 g, 0.0047 mol) was suspended in 2 M sodium hydroxide (15 ml), 20 boiled for 10 min. The pH was adjusted to 0.3 with c nc. hydrochloric acid. The solution was extracted with ethyl acetate (10 x 25 ml). The organic phase was extracted with saturated aqueous sodium chloride, dried over magnesium sulfate, and evaporated to dryness, in vacuo, to give the 25 title compound as a white solid (0.66 g, 71%). M.p. 118-121°C. Anal. for C₈H₁₀N₂O₄, found (calc.) C: 48.38 (48.49) H: 5.09 (5.09) N: 13.93 (14.14). The compound showed satisfactory ¹H and ¹³C-NMR spectra.

30 EXAMPLE 47

N-(N'-Boc-aminoethyl)-N-[(1-thyminyl)propancyl]glycine thylester.

N-(N'-Boc-aminoethyl)glycine ethyl ester (1.0 g, 0.0041 mol) was dissolved in DMF (12 ml). DhbtOH (0.73 g, 0.0045 35 mol) and 3-(1-thyminyl)-propanoic acid (0.89 g, 0.0045 mol) wer added. Methyl ne chl ride (12 ml) then was added and the mixture was cool d to 0°C on an ic bath. After addition of

DCC (1.01 g, 0.0049 mol), the mixture was stirred at 0°C for 2 h, followed by 1 h at r om temperature. The pr cipitated DCU was removed by filtration, washed with methylene chloride (25 ml), and a further amount of methylene chloride (50 ml) 5 was added to the filtrate. The organic phase was extracted with sodium hydrogen carbonate (1 volume saturated dilut d with 1 volume water, 6 x 100 ml), potassium sulfate (1 volume saturated diluted with 4 volumes water, 3 x 100 ml), and saturated aqueous sodium chloride (1 x 100 ml), dried over 10 magnesium sulfate, and evaporated to dryness, in vacuo. The solid residue was suspended in methylene chloride (15 ml), and The precipitated DCU was removed by stirred for 1h. filtration and washed with methylene chloride. The filtrate was evaporated to dryness, in vacuo, and the residue purifi d 15 by column chromatography on silica gel, eluting with a mixture of methanol and methylene chloride (gradient from 1 t 6% methanol in methylene chloride). This afforded the title compound as a white solid (1.02 g, 59%). Anal. for C10H10N2O7, found (calc.) C: 53.15 (53.51) H: 6.90 (7.09) N: 12.76 20 (13.13). The compound showed satisfactory ¹H and ¹³C-NMR spectra.

EXAMPLE 48

N-(N'-Boc-aminoethyl)-N-[(1-thyminyl)propancyl]glycine .

N-(N'-Boc-aminoethyl)-N-[(1-thyminyl)propanoyl]glycine ethyl ester (0.83 g, 0.00195 mol) was dissolved in methanol (25 ml). Sodium hydroxide (25 ml; 2 M) was added. The solution was stirred for 1 h. The methanol was removed by evaporation, in vacuo, and the pH adjusted to 2 with 4 M hydrochloric acid at 0°C. The product was isolated by filtration, washed with ether (3 x 15 ml), and dried over sicapent, in vacuo. Yield 0.769 g, 99%). M.p. 213°C (decomp.).

M no-B c-ethylenediamine (2).

t rt-Butyl-4-nitrophenyl carbonate (1) (10.0 g; 0.0418 mol) dissolved in DMF (50 ml) was added dropwise over a period 5 of 30 min to a solution of ethylenediamine (27.9 ml; 0.418 mol) and DMF (50 ml) and stirred overnight. The mixture was evaporated to dryness, in vacuo, and the resulting oil dissolved in water (250 ml). After cooling to 0°C, pH was adjusted to 3.5 with 4 M hydrochloric acid. The solution then 10 was filtered and extracted with chloroform (3x250 ml). pH was adjusted to 12 at 0°C with 2 M sodium hydroxide, and the aqueous solution extracted with methylene chloride (3x300 ml). After treatment with sat. aqueous sodium chloride (250 ml), the methylene chloride solution was dried over magnesium 15 sulfate. After filtration, the solution was evaporated t dryness, in vacuo, resulting in 4.22 g (63%) of the product (oil). H-NMR (90 MHz; CDCl₃): 61.44 (s, 9H); 2.87 (t, 2H); 3.1 (q, 2H); 5.62 (s, broad).

20 EXAMPLE 50

(N-Boc-aminoethyl)- β -alamine methyl ester, HCl.

Mono-Boc-ethylenediamine (2) (16.28 g; 0.102mol) was dissolved in acetonitrile (400 ml) and methyl acrylate (91.50 ml; 1.02 mol) was transferred to the mixture with acetonitril (200 ml). The solution was refluxed overnight under nitrog n in the dark to avoid polymerization of methyl acrylate. Aft r evaporation to dryness, in vacuo, a mixture of water and ether (200 + 200 ml) was added, and the solution was filtered and vigorously stirred. The aqueous phase was extracted one mor 100 time with ether and then freeze dried to yield a yellow solid. Recrystallization from ethyl acetate yielded 13.09 g (46%) f the title compound. M.p. 138-140°C. Anal. for C₁₁H₂₃N₂O₄Cl, found (calc.) C: 46.49 (46.72) H: 8.38 (8.20) N: 9.83 (9.91) Cl: 12.45 (12.54). H-NMR (90 MHz; DMSO-d₆): 6 1.39 (s, 9H);

N-[(1-Thyminyl)acetyl]-N'-Boc-aminoethyl- β -alanine m thylester.

 $(N-Boc-amino-ethyl)-\beta-alanine methyl ester, HCl (3)$ 5 (2.0 and 1-thyminylacetic 0.0071 mol) acid pentafluorophenyl ester (5) (2.828 g; 0.00812 mol) were dissolved in DMF (50 ml). Triethyl amine (1.12 ml; 0.00812 mol) was added and the mixture stirred overnight. addition of methylene chloride (200 ml) the organic phase was 10 extracted with aqueous sodium hydrogen carbonate (3x250 ml), half-sat. aqueous potassium hydrogen sulfate (3x250 ml), and sat. aqueous sodium chloride (250 ml) and dried over magnesium sulfate. Filtration and evaporation to dryness, in vacuo, resulted in 2.9 g (99%) yield (oil). H-NMR (250 MHz; CDCl₂): 15 due to limited rotation around the secondary amide several of the signals were doubled; δ 1.43 (s, 9H); 1.88 (s, 3H); 2.63 (t, 1H); 2.74 (t, 1H); 3.25-3.55 (4xt, 8H); 3.65 (2xt, 2H); 3.66 (s, 1.5); 3.72 (s, 1.5); 4.61 (s, 1H); 4.72 (s, 2H); 5.59 (s, 0.5H); 5.96 (s, 0.5H); 7.11 (s, 1H); 10.33 (s, 1H).

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EXAMPLE 52

$N-[(1-Thyminyl)acetyl]-N'-Boc-aminoethyl-\beta-alanine.$

N-[(1-Thyminyl)acetyl]-N'-Boc-aminoethyl-β-alanine methyl ester (3.0 g; 0.0073 mol) was dissolved in 2 M sodium 25 hydroxide (30 ml), the pH adjusted to 2 at 0°C with 4 M hydrochloric acid, and the solution stirred for 2 h. The precipitate was isolated by filtration, washed three times with cold water, and dried over sicapent, in vacuo. Yield 2.23 g (77%). M.p. 170-176°C. Anal. for C₁₇H₂₆N₄O₇, H₂O, found 30 (calc.) C: 49.49 (49.03) H: 6.31 (6.78) N: 13.84 (13.45). H-NMR (90 MHz; DMSO-d₆): δ 1.38 (s, 9H); 1.76 (s, 3H); 2.44 and 3.29 (m, 8H); 4.55 (s, 2H); 7.3 (s, 1H); 11.23 (s, 1H). FAB-MS: 399 (M+1).

 $N-[(1-(N'-Z)-cyt syl)ac tyl]-N'-B c-aminoethyl-\beta-alani ne methyl ester.$

 $(N-Boc-amino-ethyl)-\beta-alanine methyl ester, HCl (3)$ 5 (2.0 g; 0.0071 mol) and 1-(N-4-Z)-cytosylacetic acid pentafluorophenyl ester (5) (3.319 g; 0.0071 mol) were dissolved in DMF (50 ml). Triethyl amine (0.99 ml; 0.0071 mol) was added and the mixture stirred overnight. addition of methylene chloride (200 ml), the organic phase was 10 extracted with aqueous sodium hydrogen carbonate (3x250 ml), half-sat. aqueous potassium hydrogen sulfate (3x250 ml), and sat. aqueous sodium chloride (250 ml), and dried over magnesium sulfate. Filtration and evaporation to dryness, in vacuo, resulted in 3.36 g of solid compound which was 15 recrystallized from methanol. Yield 2.42 g (64%). M.p. 158-161°C. Anal. for $C_{25}H_{33}N_5O_8$, found (calc.) C: 55.19 (56.49) H: 6.19 (6.26) N: 12.86 (13.18). 1H-NMR (250 MHz; CDCl₃): due t limited rotation around the secondary amide several of the signals were doubled; & 1.43 (s, 9H); 2.57 (t, 1H); 3.60-3.23 20 (m's, 6H); 3.60 (s, 1,5H); 3.66 (s, 1.5H); 4.80 (s, 1H); 4.88 (s, 1H); 5.20 (s, 2H); 7.80-7.25 (m's, 7H). FAB-MS: 532 (M+1).

EXAMPLE 54

25 N-[(1-(M-Z)-cytosyl)acetyl]-N'-Boc-aminoethyl- β -alanine.

N-[(1--(N-4-Z)-cytosyl)acetyl]-N'-Boc-aminoethyl-\$alanine methyl ester (0.621 g; 0.0012 mol) was dissolved in
2 M sodium hydroxide (8.5 ml) and stirred for 2h.
Subsequently, pH was adjusted to 2 at 0°C with 4 M
30 hydrochloric acid and the solution stirred for 2 h. The
precipitate was isolated by filtration, washed three times
with cold water, and dried over sicapent, in vacuo. Yield
0.326 g (54%). The white solid was recrystallized from 2propanol and washed with petroleum ether. Mp.163°C (decomp.).
35 Anal. for C₂₄H₃₁N₅O₈, found (calc.) C: 49.49 (49.03) H: 6.31
(6.78) N: 13.84 (13.45). H-NMR (250 MHz; CDCl₃): due to
limited rotati n around th secondary amide several f th

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signals were doubled; & 1.40 (s, 9H); 2.57 (t, 1H); 2.65 (t, 1H); 3.60-3.32 (m's, 6H); 4.85 (s, 1H); 4.98 (s, 1H); 5.21 (s, 2H); 5.71 (s, 1H, broad); 7.99-7.25 (m's, 7H). FAB-MS: 518 (M+1).

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EXAMPLE 55

Example of a PNA-oligomer with a quanine residue

- (a) Solid-Phase Synthesis of H-[Taeg],-[Gaeg]-[Taeg],-Lys-NH,
- The protected PNA was assembled onto a Boc-Lys(C12) modified MBHA resin with a substitution of approximately 0.15 mmol/g (determined by quantitative Ninhydrin reaction). Capping of uncoupled amino groups was only carried out before the incorporation of the BocGaeg-OH monomer.
- 15 (b) Stepwise Assembly of H-[Taeg],-[Gaeg]-[Taeg],-Lys-NH; (synthetic protocol)

Synthesis was initiated on 102 mg (dry weight) of preswollen (overnight in DCM) and neutralized Boc-Lys(ClZ)-MBHA resin. The steps performed were as follows: (1) Boc-20 deprotection with TFA/DCM (1:1, v/v), 1 x 2 min and 1 x 1/2 h, 3 ml; (2) washing with DCM, 4 x 20 sec, 3 ml; washing with DMF, 2 x 20 sec, 3 ml; washing with DCM, 2 x 20 sec, 3 ml, and drain for 30 sec; (3) neutralization with DIEA/DCM (1:19 v/v), 2 x 3 min, 3 ml; (4) washing with

- with DIEA/DCM (1:19 V/V), 2 x 3 min, 3 mi; (4) washing with 25 DCM, 4 x 20 sec, 3 ml, and drain for 1 min.; (5) addition of 4 equiv. diisopropyl carbodiimide (0.06 mmol; 9.7 μl) and 4 equiv. (0.06 mmol; 24 mg) BocTaeg-OH or (0.06 mmol; 30 mg) BocGaeg-OH dissolved in 0.6 ml DCM/DMF (1:1, v/v) (final concentration of monomer 0.1 M), the coupling reaction was
- allowed to proceed for 1/2 h shaking at room temperature; (6) drain for 20 sec; (7) washing with DMF, 2 x 20 sec and 1 x 2 min, 3 ml; washing with DCM 4 x 20 sec, 3 ml; (8) neutralization with DIEA/DCM (1:19 v/v), 2 x 3 min, 3 ml; (9) washing with DCM 4 x 20 sec, 3 ml, and drain for 1 min.; (10)
- 35 qualitative Kaiser test; (11) blocking of unreacted amino groups by acetylati n with Ac₂O/pyridine/DCM (1:1:2, v/v), 1 x 1/2 h, 3 ml; and (12) washing with DCM, 4 x 20 sec, 2 x 2

min and 2 x 20 sec, 3 ml. Steps 1-12 were r p ated until the d sired sequence was obtained. All qualitative Kaiser tests were negative (straw-yellow col ur with n col ration f the beads) indicating near 100% coupling yield. The PNA-oligomer was cleaved and purified by the normal procedure. FAB-MS: 2832.11 [M+1] (calc. 2832.15)

EXAMPLE 56

Solid-Phase Synthesis of H-Taeg-Aaeg-[Taeg].-Lys-NH,.

10 (a) Stepwise Assembly of Boc-Taeg-A(2)aeg-[Taeg]:Lys(Cl2)-MBHA Resin.

About 0.3 g of wet Boc-[Taeg].-Lys(Cl2)-MBHA resin was placed in a 3 ml SPPS reaction vessel. Boc-Taeg-A(Z)aeg-[Taeg].-Lys(Cl2)-MBHA resin was assembled by in situ DCC coupling (single) of the A(Z)aeg residue utilizing 0.19 M of BocA(Z)aeg-OH together with 0.15 M DCC in 2.5 ml 50% DMF/CH₂Cl₂ and a single coupling with 0.15 M BocTaeg-OPfp in neat CH₂Cl₂ ("Synthetic Protocol 5"). The synthesis was monitored by the quantitative ninhydrin reaction, which showed about 50% incorporation of A(Z)aeg and about 96% incorporation of Ta g.

(b) Cleavage, Purification, and Identification of H-Taeg-Raeg-[Taeg],-Lys-NH2.

The protected Boc-Taeg-A(Z)aeg-[Taeg].-Lys(ClZ)-BAH resin was treated as described in Example 40c to yield about 15.6 mg of crude material upon HF cleavage of 53.1 mg dry H-Taeg-A(Z)aeg-[Taeg].-Lys(ClZ)-BHA resin. The main peak at 14.4 min accounted for less than 50% of the total absorbance. A 0.5 mg portion of the crude product was purified to give approximately 0.1 mg of H-Taeg-Aaeg-[Taeg].-Lys-NH2. For (MH+) the calculated m/z value was 2816.16 and the measured m/z value was 2816.28.

- (c) Synthetic Protocol 5
- (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, V/V), 2.5 ml, 3 x 1 min and 1 x 30 min; (2) washing with CH₂Cl₂, 2.5 ml, 6 x 1 min; (3) n utralization with DIEA/CH₂Cl₂ (1: 19, V/V), 2.5 ml, 3 x 2 min; (4) washing with CH₂Cl₂, 2.5 ml, 6 x 1 min, and drain f r 1 min; (5) 2-5 mg sampl f PNA-resin is tak n out

and dried thoroughly for a quantitative ninhydrin analysis to determine the substitution; (6) addition of 0.47 mmol (0.25 g) BocA(Z)aeg-OH dissolved in 1.25 ml DMF followed by addition of 0.47 mmol (0.1 g) DCC in 1.25 ml CH₂Cl₂ or 0.36 mmol (0.20 5 g) BocTaeg-OPfp in 2.5 ml CH₂Cl₂; the coupling reaction was allowed to proceed for a total of 20-24 hrs shaking; (7) washing with DMF, 2.5 ml, 1 x 2 min; (8) washing with CH,Cl,, 2.5 ml, 4 x 1 min; (9) neutralization with DIEA/CH,Cl, (1: 19, v/v), 2.5 ml, 2 x 2 min; (10) washing with CH₂Cl₂, 2.5 ml, 6 10 x 1 min; (11) 2-5 mg sample of protected PNA-resin is taken out and dried thoroughly for a quantitative ninhydrin analysis to determine the extent of coupling; (12) blocking of unreacted amino groups by acetylation with a 25 ml mixture of acetic anhydride/pyridine/CH₂Cl₂ (1:1:2, v/v/v) for 2 h (except 15 after the last cycle); and (13) washing with CH,Cl,, 2.5 ml, 6 x 1 min; (14) 2 x 2-5 mg samples of protected PNA-resin are taken out, neutralized with DIEA/CH₂Cl₂ (1: 19, v/v) and washed with CH,Cl, for ninhydrin analyses.

20 EXAMPLE 57

Solid-Phase Synthesis of H-[Taeg],-Aaeg-[Taeg],-Lys-NH,.

(a) Stepwise Assembly of Boc-[Taeg];-A(Z)aeg-[Taeg];-Lys(ClZ)-MBHA Resin.

About 0.5 g of wet Boc-[Taeg];-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg];-A(Z)aeg-[Taeg];-Lys(ClZ)-MBHA resin was assembled by in situ DCC coupling of both the A(Z)aeg and the Taeg residues utilising 0.15 M to 0.2 M of protected PNA monomer (free acid) together with an equivalent amount of DCC in 2 ml neat CH;Cl; 30 ("Synthetic Protocol 6"). The synthesis was monitored by the quantitative ninhydrin reaction which showed a total of about 82% incorporation of A(Z)aeg after coupling three times (the first coupling gave about 50% incorporation; a fourth HOBt-mediated coupling in 50% DMF/CH2Cl2 did not increase the total coupling yield significantly) and quantitative incorporation (single couplings) of the Taeg residues.

(b) Cleavag, Purificati n, and Id ntificati n f H[Taeg],-Aa g-[Ta g],-Lys-NH,.

The protected Boc-[Taeg],-A(Z)aeg-[Taeg],-Lys(ClZ)-BHA resin was treated as described in Example 40c to yield about 5 16.2 mg of crude material upon HF cleavage of 102.5 mg dry H-[Taeg],-A(Z)aeg-[Taeg],-Lys(ClZ)-BHA resin. A small portin of the crude product was purified. For (MH+)*, the calculated m/z value was 2050.85 and the measured m/z value was 2050.90

(c) Synthetic Protocol 6

10 (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, V/V), 2 ml, 3 \times 1 min and 1 \times 30 min; (2) washing with CH₂Cl₂, 2 ml, 6 \times 1 min; (3) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 2 ml, 3 x 2 min; (4) washing with CH2Cl2, 2 ml, 6 x 1 min, and drain for 1 min; (5) 2-5 mg sample of PNA-resin was taken out and 15 dried thoroughly for a quantitative ninhydrin analysis to determine the substitution; (6) addition of 0.44 mmol (0.23 g) BocA(Z)aeg-OH dissolved in 1.5 ml CH₂Cl₂ followed by addition of 0.44 mmol (0.09 g) DCC in 0.5 ml CH2Cl2 or 0.33 mmol (0.13 g) BocTaeg-OH in 1.5 ml CH,Cl, followed by additi n 20 of 0.33 mmol (0.07 g) DCC in 0.5 ml CH₂Cl₂; the coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking; (7) washing with DMF, 2 ml, 1 x 2 min; (8) washing with CH₂Cl₂, 2 ml, 4 x 1 min; (9) neutralization with DIEA/CH,Cl2 (1: 19, v/v), 2 ml, 2 x 2 min; (10) washing with 25 CH₂Cl₂, 2 ml, 6 x 1 min; (11) 2-5 mg sample of protected PNAresin is taken out and dried thoroughly for a quantitative ninhydrin analysis to determine the extent of coupling; (12) blocking of unreacted amino groups by acetylation with a 25 ml mixture of acetic anhydride/pyridine/CH2Cl2 (1:1:2, V/V/V) 30 for 2 h (except after the last cycle); (13) washing with CH_1Cl_2 , 2 ml, 6 x 1 min; and (14) 2 x 2-5 mg samples f protected PNA-resin were taken out, neutralized with DIEA/CH,Cl, (1: 19, v/v) and washed with CH,Cl, for ninhydrin analyses.

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EXAMPLE 58

The PNA-oligomer H-T4C2TCT-LysNH₂ was prepared as described in Example 93. Hybridization experiments with this sequence should resolve the issue of orientation, since it is truly asymmetrical. Such experiments should also resolve the issues of pH-dependency of the Tm, and the stoichiometry of complexes formed.

Hybridization experiments with the PNA-oligomer H- $T_{\ell}C_{2}TCTC$ -LysNH $_{2}$ were performed as follows:

الكامانية المساعر		
Row	Hybridized With	
1	5'-(dA),(dG),(dA)(dG)(dA)(dG)	7

i	NOW	Hybridized with) PA	1,444	
	1	5'-(dA),(dG),(dA) (dG) (dA) (dG)	7.2	55.5	2:1
	2	5'-(dA),(dG),(dA)(dG)(dA)(dG)	9.0	26.0	2:1
	3	5'-(dA),(dG),(dA)(dG)(dA)(dG)	5.0	88:5	2:1
15	4	5'-(dG)(dA)(dG)(dA)(dG),(dA),	7.2	38.0	2:1
	5	5'-(dG)(dA)(dG)(dA)(dG),(dA),	9.0	31.5	-
	6	5'-(dG)(dA)(dG)(dA)(dG),(dA),	5.0	52.5	-
	7	5'-(dA),(dG)(dT)(dA)(dG)(dA)(dG)	7.2	39.0	-
	8	5'-(dA),(dG)(dT)(dA)(dG)(dA)(dG)	9.0	<20	-
20	9	5'-(dA),(dG)(dT)(dA)(dG)(dA)(dG)	5.0	51.5	
	10	5'-(dA),(dG),(dT)(dG)(dA)(dG)	7.2	31.5	
	11	5'-(dA),(dG),(dT)(dG)(dA)(dG)	5.0	50.5	-
	12	5'-(dG)(dA)(dG)(dA)dT)(dG)(dA),	7.2	24.5	-
	13	5'-(dG)(dA)(dG)(dA)dT)(dG)(dA),	9.0	<20	_
25	14	5'-(dG)(dA)(dG)(dA)dT)(dG)(dA),	5.0	57.0	-
1	15	5'-(dG)(dA)(dG)(dT)(dG),(dA),	7.2	25.0	_
	16	5'-(dG)(dA)(dG)(dT)(dG),(dA),	5.0	39.5	_
				F2 0	

= stoichiometry determined by UV-mixing curves

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- = not determined

These results show that a truly mixed sequence gave rise to well defined melting curves. The PNA-oligomers can actually bind in both orientations (compare row 1 and 4), 35 although there is preference for the N-terminal/5'-

orientation. Introducing a single mismatch pposit either T or C caus d a lowering f T_a by mor than 16°C at pH 7.2; at pH 5.0 the T_a-value was lowered more than 27°C. This shows that there is a very high degree a sequence-selectivity which should be a general feature for all PNA C/T sequences.

As indicated above, there is a very strong pH-dependency for the T_m-value, indicating that Heogsteen basepairing is important for the formation of hybrids. Therefore, it is not surprising that the stoichiometry was 10 found to be 2:1.

The lack of symmetry in the sequence and the very large lowering of T_m when mismatches are present show that th Watson-Crick strand and the Hoogsteen strand are parallel when bound to complementary DNA. This is true for both of the orientations, i.e., 5'/N-terminal and 3'/N-terminal.

EXAMPLE 59

The results of hybridization experiments with H-T,GT,LysNH, to were performed as follows:

20	Row	Deoxyoligonucleotide	Tm
	1,	5'-(dA)5(dA)(dA)4-3'	55.0
	2	5'-(dA)5(dG)(dA)4-3'	47.0
	3	5'-(dA)5(dG)(dA)4-3'	56.5
	4	5'-(dA)5(dT)(dA)4-3'	46.5
25	5	5'-(dA)4(dG)(dA)5-3'	48.5
	6	5'-(dA)4(dC)(dA)5-3'	55.5
	7	5'-(dA)4(dT)(dA)5-3'	47.0

As shown by comparing rows 1, 3, and 6 with rows 2, 4, 30 5, and 7, G can in this mode discriminate between C/A and G/T in the DNA-strand, i.e., sequence discrimination is observed. The complex in row 3 was furthermore determined to be 2 PNA: 1 DNA complex by UV-mixing curves.

EXAMPLE 60

The masses of some synthesized PNA-oligomers, as determined by FAB mass spectrometry, are as follows:

	Sequence	CALC.	POUND
5	H-T_C,TCTC-LysNH,	2747.15	2746.78
	H-T ₅ GT ₄ -LysnH ₂	2832.15	2832.11
	H-T,-LysNH,	2008.84	2540.84
10	H-To-LysnH,	2541.04	2540.84
	H-T ₁₀ -LysnH ₂	2807.14	2806.69
	H-T,CT,-LysnH,	2259.94	2259.18
	$H-T_3(L-alaT)T_1-LysnH_2$	2287.95	2288.60
	H-T ₄ (Ac) T ₅ -LysNH ₂	2683.12	2683.09

EXAMPLE 61

15 Hybridization data for a PNA-oligomer with a single unit with an extended backbone (the β -alanine modification) is as follows:

	PNA	DNA	T	
	H-T _{in} -LysnH,	(dA) ₁₀	73°C	
20	H-T ₄ (βT)T ₅ -LysNH,	(dA) 10	57°C	
	$H-T_L(\beta T)T_S-LysnH_S$	(dA) (dG) (dA),	47°C	
	H-T, (BT) T,-LysnH,	(Ab) (Tb) (Ab)	49°C	
	H-T ₄ (βT)T ₅ -LysNH,	(dA) (dT) (dA),	47°C	

25 Although the melting temperature decreases, the data demonstrates that base specific recognition is retained.

An exampl with a "no base" substitution.

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	PNA	DNA	T
	H-T ₁₀ -LysNH ₂	(dA) 10	73°C
20	H-T, (Ac) T,-LysnH,	(dA),0	49°C
	H-T, (Ac) T,-LysnH,	(dA), (dG) (dA) 5	37°C
	H-T, (Ac) T,-LysnH,	$(dA)_{\ell}(dC)(dA)^{5}$	41°C
	H-T, (Ac) T,-LysnH,	(da), (dt) (da) 5	41°C
İ	H-T, (Ac) T,-LysnH,	(dA), (dG) (dA) ⁴	36°C
25	H-T, (Ac) T,-LysnH,	(da),(dc) (da)4	40°C
	H-T, (Ac) T,-LysnH,	(dA),(dT)(dA) ⁴	40°C

EXAMPLE 63

Iodination Procedure

A 5 μg portion of Tyr-PNA-T₁₀-Lys-NH₂ is dissolved in 40 μl 100 mM Na-phosphate, pH 7.0, and 1 mCi Na¹²⁵I and 2 μl chloramine-T (50 mM in CH₃CN) are added. The solution is left at 20°C for 10 min and then passed through a 0.5 + 5 cm Sephadex G10 column. The first 2 fractions (100 μl each) containing radioactivity are collected and purified by HPLC: reversed phase C-18 using a 0-60% CH₃CN gradient in 0.1% CF₃COOH in H₂O. The ¹²⁵I-PNA elutes right after the PNA p ak. The solvent is removed under reduced pressure.

Binding of PNAs- $T_{10}/T_{\phi}C/T_{g}C_{2}$ to double stranded DNA targets $\lambda_{10}/\lambda_{\phi}G/\lambda_{g}G_{2}$ (Pigure 20).

A mixture of 200 cps ³²P-labeled EcoRI-PvuII fragment 5 (the large fragment labeled at the 3'-end of the EcoRI site) of the indicated plasmid, 0.5 μg carrier calf thymus DNA, and 300 ng PNA in 100 μl buffer (200 mM NaCl, 50 mM Na-acetate, pH 4.5, 1 mM ZnSO₄) was incubated at 37°C for 120 min. A 50 unit portion of nuclease S₁ was added and incubated at 20°C for 5 min. The reaction was stopped by addition of 3 μl 0.5 M EDTA and the DNA was precipitated by addition of 250 μl 2% potassium acetate in ethanol. The DNA was analyzed by electrophoresis in 10% polyacrylamide sequencing gels and the radiolabeled DNA bands visualized by autoradiography.

The target plasmids were prepared by cloning of the appropriate oligonucleotides into pUC19. Target A₁₀: oligonucleotides GATCCA₁₀G & GATCCT₁₀G cloned into the BamHI site (plasmid designated pT10). Target A₅GA₄: oligonucleotides TCGACT₄CT₅G & TCGACA₅GA₄G cloned into the SalI site (plasmid pT9C). Target A₂GA₂GA₄: oligonucleotides GA₂GA₂GA₄TGCA & GT₄CT₂CT₂CTGCA into the PstI site (plasmid pT8C2). The positions of the targets in the gel are indicated by bars to the left. A/G is an A+G sequence ladder of target F10.

25 EXAMPLE 65

Inhibition of restriction enzyme cleavage by PNA (Figure 23).

A 2 μ g portion of plasmid pT10 was mixed with the indicated amount of PNA-T₁₀ in 20 μ l TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 7.4) and incubated at 37°C for 120 min. 2 30 μ l 10 × buffer (10 mM Tris-HCl, pH 7.5, 10 mM, MgCl₂, 50 mM NaCl, 1 mM DTT). PvuII (2 units) and BamHI (2 units) were added and the incubation was continued for 60 min. The DNA was analyzed by gel electrophoresis in 5% polyacrylamide and the DNA was visualized by ethidium bromide staining.

Kinetics of PNA- T_{10} - dsDNA strand displa ement c mplex f rmati n (Figure 21).

A mixture of 200 cps ³²P-labeled EcoRI-PvuII fragment of pT10 (the large fragment labeled at the 3'-end of the EcoRI site), 0.5 μg carrier calf thymus DNA, and 300 ng of PNA-T₁₀-LysNH₂ in 100 μl buffer (200 mM NaCl, 50 mM Na-acetate, pH 4.5, 1 mM ZnSO₄) were incubated at 37°C. At the times indicated, 50 U of S₁ nuclease was added to each of 7 samples and incubation was continued for 5 min at 20°C. The DNA was then precipitated by addition of 250 μl 2% K-acetate in ethanol and analyzed by electrophoresis in a 10% polyacrylamide sequencing gel. The amount of strand displacement complex was calculated from the intensity of the S₁-cleavage at the target sequence, as measured by densitometric scanning of autoradiographs.

EXAMPLE 67

Stability of PNA-dsDNA complexes (Figure 22).

A mixture of 200 cps ³²P-pT10 fragment, 0.5 μg calf thymus DNA and 300 ng of the desired PNA (either T₁₀-LysNH₂, T₈-LysNH₂ or T₆-LysNH₂) was incubated in 100 μl 200 mM NaCl, 50 mM Na-acetate, pH 4.5, 1 mM ZnSO₄ for 60 min at 37°C. A 2 μg portion of oligonucleotide GATCCA₁₀G was added and each sampl was heated for 10 min at the temperature indicated, cooled in ice for 10 min and warmed to 20°C. A 50 U portion of S₁ nuclease was added and the samples treated and analyzed and

30 EXAMPLE 68

Inhibition of Transcription by PNA

the results quantified.

A mixture of 100 ng plasmid DNA (cleaved with restriction enzyme PvuII (see below) and 100 ng of PNA in 15 μ l 10 mM Tris-HCl, 1 mM EDTA, pH 7.4 was incubated at 37°C f r 35 60 min. Subsequently, 4 μ l 5 × concentrated buffer (0.2 M Tris-HCl (pH 8.0), 40 mM MgCl₂, 10 mM sp rmidine, 125 mM NaCl) were mixed with 1 μ l NTP-mix (10 mM ATP, 10 mM CTP, 10 mM GTP,

1 mM UTP, 0.1 μ Ci/ μ l ³²P-UTP, 5 mM DTT, 2 μ g/ml tRNA, 1 μ g/ml heparin) and 3 units RNA polymerase. Incubation was continued for 10 min at 37°C. The RNA was then pr cipitated by addition of 60 \(\mu \)l 2% postassium acetate in 96% ethanol at -20°C and 5 analyzed by electrophoresis in 8% polyacrylamide sequencing gels. RNA transcripts were visualized by autoradiography. The following plasmids were used: pT8C2-KS/pA8G2-KS: oligonucleotides GA,GA,GA,GTGAC & GT,CT,CT,CTGCA cloned into the pBluescript-KS; of pT10-KS/pA10-KS 10 orientations of the insert were obtained). pT10UV5: oligonucleotides GATCCA, G & GATCCT, G cloned into the BamHI site of a pUC18 derivative in which the lac UV5 promoter had been cloned into the EcoRI site (Jeppesen, t al., Nucleic Acids Res., 1988, 16, 9545).

Using T₃-RNA polymerase, transcription elongation arrest was obtained with PNA-T₈C₂-LysNH₂ and the pA8G2-KS plasmid having the PNA recognition sequence on the template strand, but not with pT8C2-KS having the PNA recogniti n sequence on the non-template strand. Similar results wer obtained with PNA-T10-LysNH₂ and the plasmids pA10-KS and pT10-KS. (see, Pigure 25) Using E.coli RNA polymerase and the pT10UV5 plasmid (A₁₀-sequence on the template strand) transcription elongation arrest was obtained with PNA-T₁₀-LysNH₂.

25

EXAMPLE 69

Biological stability of PNA

A mixture of PNA-T₅ (10 μg) and a control, "normal" peptide (10 μg) in 40 μl 50 mM Tris-HCl, pH 7.4 was treated 30 with varying amounts of peptidase from porcine intestinal mucosa or protease from Streptomyces caespitosus for 10 min at 37°C. The amount of PNA and peptide was determined by HPLC analysis (reversed phase C-18 column: 0-60% acetonitrile, 0.1% trifluoroacetic acid).

At peptidase/protease concentrations where complete d gradation of th peptide was bs rved (n HPLC peak) the PNA was still intact.

Inhibiti n f Gene Expressi n

A preferred assay to test the ability of peptid nucleic acids to inhibit expression of the E2 mRNA 5 papillomavirus is based on the well-documented transactivati n properties of E2. Spalholtz, et al., J. Virol., 1987, 61, 2128-2137. A reporter plasmid (E2RECAT) was constructed t contain the E2 responsive element, which functions as an E2 E2RECAT also contains the SV40 early dependent enhancer. 10 promoter, an early polyadenylation signal, and chloramphenicol acetyl transferase gene (CAT). Within the context of this plasmid, CAT expression is dependent upon expression of E2. The dependence of CAT expression on th presence of E2 has been tested by transfection of this plasmid 15 into C127 cells transformed by BPV-1, uninfected C127 cells and C127 cells cotransfected with E2RECAT and an E2 expression vector.

A. Inhibition of BPV-1 E2 Expression

BPV-1 transformed C127 cells are plated in 12 well 20 plates. Twenty four hours prior to transfection with E2RE1, cells are pretreated by addition of antisense PNAs to the growth medium at final concentrations of 5, 15 and 30 mM. Th next day cells are transfected with 10 μg of E2RE1CAT by calcium phosphate precipitation. Ten micrograms of E2RE1CAT 25 and 10 μg of carrier DNA (PUC 19) are mixed with 62 μl of 2 M CaCl₂ in a final volume of 250 μ l of H₂0, followed by addition of 250 μ l of 2X HBSP (1.5 mM Na₂PO₂. 10 mM KCl, 280 mM NaCl, 12 mM glucose and 50 mM HEPES, pH 7.0) and incubated at room temperature for 30 minutes. One hundred microliters 30 of this solution is added to each test well and allowed t incubate for 4 hours at 37°C. After incubation, cells ar glycerol shocked for 1 minute at room temperature with 15% glycerol in 0.75 mm Na₂PO₂, 5 mm KCl, 140 mM NaCl, 6 mM glucose and 25 mM HEPES, pH 7.0. After shocking, cells are 35 washed 2 times with serum free DMEM and refed with DMEM s rum and fetal bovine containing 10% lig nucle tide at th riginal c nc ntrati n. F rty ight

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hours after transfection cells are harvested and assayed for CAT activity.

For determination of CAT activity, cells are washed 2 times with phosphate buffered saline and collected by 5 scraping. Cells are resuspended in 100 μl of 250 mM Tris-HCl, pH 8.0 and disrupted by freeze-thawing 3 times. Twenty four microliters of cell extract is used for each assay. For ach assay the following are mixed together in an 1.5 ml Eppendorf tube and incubated at 37°C for one hour: 25 μ l of cell 10 extract, 5 μ l of 4 mM acetyl coenzyme A, 18 μ l H,O and 1 μ l 14C-chloramphenicol, 40-60 mCi/mM. After incubation, chloramphenicol (acetylated and nonacetylated forms) is extracted with ethyl acetate and evaporated to dryness. Samples are resuspended in 25 μ l of ethyl acetate, spotted 15 onto a TLC plate and chromatographed in chloroform:methanol (19:1). Chromatographs are analyzed by autoradiography. Spots corresponding to acetylated and nonacetylated 14Cchloramphenicol are excised from the TLC plate and counted by liquid scintillation for quantitation of CAT activity. 20 Peptide nucleic acids that depress CAT activity in a dose dependent fashion are considered positives.

B. Inhibition of HPV E2 Expression

The assay for inhibition of human papillomavirus (HPV)
E2 by peptide nucleic acids is essentially the same as that
25 for BPV-1 E2. For HPV assays appropriate HPVs are cotransfected into either CV-1 or A431 cells with PSV2NEO using
the calcium phosphate method described above. Cells which
take up DNA are selected for by culturing in media containing
the antibiotic G418. G418-resistant cells are then analyz d
30 for HPV DNA and RNA. Cells expressing E2 are used as target
cells for antisense studies. For each PNA, cells are
pretreated as above, transfected with E2RE1CAT, and analyzed
for CAT activity as above. Peptide nucleic acids are
considered to have a positive effect if they can depress CAT
35 activity in a dose dependent fashion.

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EXAMPLE 71

Synthesis f PNA 15-mer C ntaining F ur Naturally Occurring Nucleobases; H-[Taeg]-[Aaeg]-[Gaeg]-[Taeg]-[Taeg]-[Aaeg]-[Taeg]-[Caeg]-[Taeg]-[Caeg]-[Taeg]-[Taeg]-[Caeg]-[Caeg]-[Taeg]-[Caeg]-[Taeg]-[Caeg]-[Taeg]-[Caeg]-[Taeg]-[Caeg]-[Taeg]-[Caeg]-[Caeg]-[Taeg]-[Caeg]-[Caeg]-[Taeg]-[Caeg]-[Caeg]-[Taeg]-[Caeg

The protected PNA was assembled onto a Boc-Lys(ClZ) modified MBHA resin with a substitution of approximately 0.145 mmol/g. Capping of uncoupled amino groups was only carried out before the incorporation of the BocGaeg-OH monomer.

Synthesis was initiated on 100 mg (dry weight) f neutralised Boc-Lys(ClA)-MBHA resin that had been preswollen overnight in DCM. The incorporation of the monomers followed the protocol of Example 32, except at step 5 for th incorporation of the BocAaeg-OH monomer. Step 5 for the 15 present synthesis involved addition of 4 equiv. diisopropyl carbodimide (0.06 ml; 9.7 μl) and 4 equiv. BocAaeg-OH (0.06 mmol; 32 mg) dissolved in 0.6 ml DCM/DMF (1:1, v/v) (final concentration of monomer 0.1M). The coupling reaction was allowed to proceed for 1 x 15 min and 1 x 60 min. 20 (recoupling).

All qualitative Kaiser tests were negative (strawyellow color with no coloration of the beads). The PNAoligomer was cleaved and purified by the standard procedure. FAB-MS average mass found(calc.) (M+H) 4145.1 (4146.1).

EXAMPLE 72
Hybridization of H-TAGTTATCTCTATCT-LysNH₂

DNA -target	Eq	TR
5'3'	5	60.5
5'3'	7.2	43.0
5'3'	9	38.5
3'5'	5	64.5/49.0
3'5'	7.2	53.5
3'5'	9	51.5

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The fact that there is alm st no 1 ss in Tm in going from pH 7.2 to 9.0 indicates that Hoogsteen bas pairing is not involved. The increas in Tm in going from 7.2 to 5 is large for the parallel orientation and is probably due to the 5 formation of a 2:1 complex. It is believed that the most favorable orientation in the Watson-Crick binding motif is the 3'/N-orientation and that in the Hoogsteen motif the 5'/N-orientation is the most stable. Thus, it may be the case that the most stable complex is with the two PNA's strands anti 10 parallel.

There is apparently a very strong preference for a parallel orientation of the Hoogsteen strand. This seems t explain why even at pH 9 a 2:1 complex is seen with the 5'/N-orientation. Furthermore, it explains the small loss in going 15 from pH 7.2 to 9 in the 3'/N, as this is probably a 1:1 complex.

EXAMPLE 73

Solid-Phase Synthesis of H-[Taeg],-Aaeg-Taeg-Caeg-Aaeg-Taeg-20 Caeg-Taeg-Caeg-Lys-NH2.

(a) Stepwise Assembly of Boc-[Taeg]2-A(3)aeg-Ta g-C(2)aeg-A(3)aeg-Taeg-C(3)aeg-Lys(Cl2)-MBHA Resin.

About 1 g of wet Boc-Lys(Cl2)-MBHA (0.28 mmol Lys/g)
25 resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]2A(Z)aeg-Taeg-C(Z)aeg-A(Z)aeg-Taeg-C(Z)aeg-Taeg-C(Z)aegLys(ClZ)-MBHA resin was assembled by in situ DCC coupling f
the five first residues utilizing 0.16 M of BocC[Z]-OH,
BocTaeg-OH or BocA(Z)aeg-OH, together with 0.16 M DCC in 2.0
30 ml 50% DMF/CH2Cl2 ("Synthetic Protocol 9") and by analogous
in situ DIC coupling of the five last residues ("Synthetic
Protocol 10"). Each coupling reaction was allowed to proceed
for a total of 20-24 hrs with shaking. The synthesis was
monitored by the ninhydrin reaction, which showed nearly
35 quantitative incorporation of all residues except of the first
A(Z)a g residue, which had to be coupled twice. The total

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coupling yield was about 96% (first coupling, about 89% efficiency).

(b) Cleavage, Purificati n, and Identification f H-[Taeg]2-haeg-Taeg-Caeg-haeg-Taeg-Caeg-Taeg-Caeg-Lys-NH2.

The protected Boc-[Taeg]2-A(Z)aeg-Taeg-C(Z)aeg-A(Z)aeg-Taeg-C(Z)aeg-Taeg-C(Z)aeg-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 53.4 mg of crude material upon HF cleavage of 166.1 mg dry Boc-[Taeg]2-A(Z)a g-Taeg-C(Z) aeg-A(Z) aeg-Taeg-C(Z) aeg-Lys(ClZ) -MBHA 10 resin. The crude product (53.4 mg) was purified to give 18.3 mg of H-[Taeg]2-Aaeg-Taeg-Caeg-Aaeg-Taeg-Caeg-Lys-NH2. For (M+H)+, the calculated m/z value = 2780.17 and the measured m/z value = 2780.07.

15 EXAMPLE 74

Hybridization properties of H-TTA TCA TCT C-Lys-NH2.

The title compound hybridized with the following oligonucleotides:

	Oligodeoxynucleotide	На	Tm(°C)
20	5'-AAT AGT AGT G-3	5	31.5†
I	5'-ATT AGT AGT G-3'	7.2	28.5†
I	5'-AAT AGT AGT G-3"	9	28.0†
	5'-GTG ATG ATA A-3'	7.2	30.5
	5'-GTG ATG ATA A-3'	9	28.0
:5	†Low hypochromicity		

Synthesis of a PNA With Tw Parallel Strings Tied Together

A 375 mg portion of MBHA resin (loading 0.6 mmol/g) was 15 allowed to swell over night in dichloromethane (DCM). After an hour in DMF/DCM, the resin was neutralized by washing 2 times with 5% diisopropylethylamine in DCM (2 min.), followed by washing with DCM (2ml; 6 x 1 min.) N, N'-di-Boc-aminoethyl glycine (41,9 mg; 0,132 mmol) disolved in 2 ml DMF was added 20 to the resin, followed by DCC (64,9 mg; 0,315 mmol) dissolved in 1 ml of DCM. After 2.5 hours, the resin was washed with DMF 3 times (1 min.) and once with DCM (1 min.). unreacted amino groups were then capped by treatment with acetic anhydride/DCM/pyridine (1 ml\2 ml\2 ml) for 72 hours. 25 After washing with DCM (2 ml; 4 x 1 min), a Kaiser test show d no amino groups were present. The resin was deprotected and washed as described above. This was followed by reaction with 6-(Bocamino)-hexanoic acid DHBT ester (255.8 mg; 67 mmol) dissolved in DMF/DCM 1:1 (4 ml) overnight. After washing and 30 neutraliation, a Kaiser test and an isatin test were performed. Both were negative. After capping, the elongenation of the PNA-chains was performed according to standard procedures for DCC couplings. All Kaiser tests performed after the coupling reactions were negative (Yellow). 35 Qualitative Kaiser tests were done after deprotection of PNA units number 1, 2, 4, and 6. Each test was blue. oligomers were cleaved and purified by standard procedures.

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The amount of monom r and DCC used for each c upling was as follows (total v lume 4.5 ml):

Coup	ling	Monome	r(T)	DCC	
1	•	173 1	ng .	95 mg	
2		176	ng	101 mg	,
3	•	174	ng	97 mg	
		174	ng	103 mg	1
5	•	178	ng	97 mg	
6		173	mg	99 mg	
7	•	174	ng	95 mg	
8		175	ng	96 mg	

10

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15 For the PNA having the Structure (70) where $R_{70} = T_6$, there was 24.5 mg of crude product, which resulted in 6.9 mg. after purification. For the PNA where $R_1 = T_8$, there was 28.8 mg of crude product, which resulted in 2.8 mg. after purification. The products had a high tendency of aggregation, as indicated by a complex HPLC chromatogram after a few hours at room temperature in concentration abov 1 mg/ml. The PNA- $(T_6)_2$ and PNA- $(T_8)_2$ were hybridised to $(dA)_6$ and $(dA)_8$, respectively, with recorded Tm of 42°C and 59°C, respectively.

25

EXAMPLE 76

Solid-Phase Synthesis of H-[Taeq],-Lys(Cl2)-MBHA Resin

The PNA oligomer was assembled onto 500 mg (dry weight) of MBHA resin that had been preswollen overnight in DCM. The 30 resin was initially substituted with approximately 0.15 mmol/g Boc-Lys(ClZ) as determined by quantitative ninhydrin reactin. The stepwise synthesis of the oligomer followed the synth tic protocol described in Example 32 employing 0.077 g (0.2 mmol) BocTaeg-OH and 31.3 μ l (0.2 mmol) diisopropyl carbodiimid in 35 2.0 ml 50% DMF/CH₂Cl₂ in each coupling. Capping f uncoupled amino groups was carried out before depretectin in ach step. All qualitative Kaiser tests were negative indicating near 100% coupling yield.

S lid-Phase Synthesis of H-[Taeg],-[apgT]-[Taeg],Lys-NH,

Synthesis was initiated on approximately 1/4 of the wet H-[Taeg]5-Lys(Cl2)-MBHA resin from Example 76. In situ diisopropyl carbodiimide (DIC) couplings of both Boc-(apgT)-OH and BocTaeg-OH were carried out in 1.2 ml 50% DMF/CH₂Cl₂ using 0.048 g (0.12 mmol) and 0.046 g (0.12 mmol) monomer, respectively, and 18.7 µl (0.12 mmol) diisopropyl carbodiimide in each coupling. All qualitative Kaiser tests were negative, indicating near 100% coupling yield. The PNA oligomer was cleaved and purified by standard procedures. For (M+H)+, the calculated m/z value was 2820.92.

15 EXAMPLE 78

Solid-Phase Synthesis of H-[Taeg],-[proT]-[Taeg],-Lys-NH, -

Synthesis was initiated on approximately 1/4 of the wet H-[Taeg],-Lys(Cl2)-MBHA resin from Example 76. In situ diisopropyl carbodiimide couplings of BocTaeg-OH were carried out in 1.2 ml 50% DMF/CH,Cl, using 0.046 g (0.12 mmol) monomer and 18.7 µl (0.12 mmol) diisopropyl carbodiimide in each coupling. Due to solubility problems, Boc-(proT)-OH 0.048 g (0.12 mmol) was suspended in 2.5 ml 50% DMF/DMSO prior to coupling, the suspension filtered, and approximately 2 ml of the filtrate used in the overnight coupling. All qualitative Kaiser tests were negative, indicating near 100% coupling yield. The PNA oligomer was cleaved and purified by standard procedures.

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EXAMPLE 79

Hybridizati n pr perties f H-[Ta g],-[pr T]-[Taeg]5Lys-NH,

Oligodeoxynucleotide	Tm(°C)
5'-AAA AAA AAA A	53.5
5-'AAA AGA AAA A	44.0
5'-AAA AAG AAA A	43.5
5'-AAA ACA AAA A	46.5
5'-AAA AAC AAA A	46.5
5'-AAA ATA AAA A	46.5
5'-AAA AAT AAA A	46.0

10

5

EXAMPLE 80 Solid-Phase Synthesis of H-[Taeg],-[bC]-[Taeg],-Lys-NH,

The PNA oligomer was assembled onto 100 mg (dry weight)
MBHA resin that had been preswollen overnight in DCM. The
resin was initially substituted with approximately 0.25 mmol/g
Boc-Lys(Cl2) as determined by quantitative ninhydrin reaction.
The stepwise synthesis of the oligomer followed synthetic
Protocol 9 employing 0.023 g (0.06 mmol) BocTaeg-OH, 0.062 g
(0.12 mmol) BocbC(Z)-OH and 0.012 g (0.06 mmol) DCC in 1.2 ml
50% DMF/CH₂Cl₂ in each coupling. Capping of uncoupled amino
groups was carried out before deprotection in each step. All
qualitative Kaiser tests were negative, indicating near 100%
coupling yield. The PNA-oligomer was cleaved and purified by
standard procedures.

EXAMPLE 81

Hybridization pr perties f H-T,bCT,-Lys-NH,

5	Oligodeoxynucleotide	Tm(°C)
	5'-AAA AAA AAA A	43.5
	5-'AAA AGA AAA A	58.0
	5'-AAA AAG AAA A	60.0
	5'-AAA ACA AAA A	34.5
10	5'-AAA AAC AAA A	34.5
	5'-AAA ATA AAA A	34.0
	5'-AAA AAT AAA A	36.0

15 Stepwise Assembly of H-[Taeg]-[Taeg]-[Taeg]-[Taeg]-[Aaeg][Taeg]-[Taeg]-[Taeg]-[Taeg]-LYS-NH,.

Synthesis was initiated on a Boc-[Taeg],-Lys(Cl2)-MBHA resin (from example 76) that had been preswollen overnight in DCM. The resin resembled approximately 100 mg (dry Weight) 20 of Boc-Lys(Cl2)-MBHA resin (loading 0.15 mmol/g). The incorporation of the monomers followed the protocol of example 55, except for step 5 (incorporation of the BocA(Z)aeg-OH monomer). New step 5 (incorporation of A(Z)aeg) involved addition of 4 equiv. diisopropyl carbodiimide (0.06 mmol; 9.7 25 µl) and 4 equiv. BocA(Z)aeg-OH (0.06 mmol; 32 mg) dissolved in 0.6 ml DCM/DMF (1:1, v/v) (final concentration of monomer 0.1 M). The coupling reaction was allowed to proceed for 1 x 15 min. and 1 x 60 min. (recoupling).

Capping of uncoupled amino groups was only carried ut 30 before the incorporation of the BocA(Z)aeg-OH monomer. The coupling reaction was monitored by qualitative ninhydrin reaction (Kaiser test). All qualitative Kaiser tests were negative (straw-yellow color with no coloration of the beads). The PNA oligomer was cleaved and purified by standard procedures.

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EXAMPLE 84

Hybridizati n properties f H-T,AT,-LysNH,

Oligodeoxynucleotide	Tm(°C)
5'-AAA AAA AAA A -	59.5
5-'AAA AGA AAA A	45.0
5'-AAA AAG AAA A	45.5
5'-AAA ACA AAA A	48.0
5'-AAA AAC AAA A	48.0
5'-AAA ATA AAA A	52.0
5'-AAA AAT AAA A	52.5

10

5

EXAMPLE 85

Stepwise Assembly of H-[Taeg]-[Taeg]-[Taeg]-[Gaeg]-15 [Gaeg]-[Taeg]-[Gaeg]-[Taeg]-[Gaeg]-LYS-NH2.

The protected PNA was assembled onto a Boc-Lys(ClZ) modified MBHA resin with a substitution of 0.15 mmol/g. The incorporation of the monomers followed the protocol of example 32, except that the capping step 11 and the washing step 12 20 were omitted. After the incorporation and deprotection of th first, second, and fourth G(Bzl)aeg-monomer there were s me difficulties getting the resin to swell properly. Three hours of shaking in neat DCM gave acceptable swelling. incorporation of residues Taeg-4, G(Bzl)aeg-6, and Taeg-7 t 25 Taeg-10, recoupling was necessary to obtain near quantitative coupling yields. Taeg, (2 x in 50% DMF/DCM), Gaeg, (2 x in 50% DMF/DCM), Taeg, (2 x in 50% DMF/DCM, 1 x in 50% NMP/DCM and 1 x in neat DCM), Taeg₈ (1 x in 50% DMF/DCM and 2 x in neat DCM), Taeg, (2 x in 50% DMF/DCM), Taeg, (2 x in 50% DMF/DCM). 30 All qualitative Kaiser tests were negative (straw-yellow color The PNA oligomer was with no coloration of the beads). cleaved and purified by standard procedures

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EXAMPLE 86

Hybridization pr perties f crude (appr x. 50%) H-T₁G₂TGTG-LysNH.

Oligodeoxynucleotide	Tm
5'-A4C2ACAC	38
5'-CACAC2A4	55

EXAMPLE 87 .

- 10 Large Scale Solid-Phase Synthesis of H-[Taeg],-Lys-NH2, H[Taeg],-Lys-NH2, H-[Taeg],-Lys-NH2, H-[Taeg],-Lys-NH2, and H[Taeg],-Lys-NH2.
 - (a) Stepwise Assembly of Boc-[Taeg] $_{10}$ -Lys(ClZ)-MBHA Resin and Shorter Fragments.
- About 9 g of wet Boc-[Taeg],-Lys(Cl2)-MBHA (s e, 15 Example 19b) resin was placed in a 60 ml SPPS reaction vessel. Boc-[Taeg],-Lys(Cl2)-MBHA resin was assembled by single coupling of both residues with 0.15 M of BocTaeg-OPfp in 10 ml neat CH₂Cl₂ ("Synthetic Protocol 8"). Both coupling 20 reactions were allowed to proceed overnight. The synthesis was monitored by the ninhydrin reaction, which showed cl se to quantitative incorporation of both residues. deprotection of the N-terminal Boc group, about 4.5 g of H-[Taeg]:-Lys(Cl2)-MBHA was placed in a 20 ml SPPS reacti n 25 vessel and elongated to Boc-[Taeg].-Lys(ClZ)-MBHA by single in situ DCC coupling of all residues (close to quantitativ , except for residue number eight) overnight with 0.2 M of BocTaeg-OH together with 0.2 M DCC in 7.5 ml neat CH,Cl, ("Synthetic Protocol 9"). Before coupling of Taeg residues 30 number seven and eight, respectively, small portions of H-[Taeg],-Lys(ClZ)-MBHA and H-[Taeg],-Lys(ClZ)-MBHA, respectively, were taken out for HF cleavage.

Taeg residue number eight was coupled twice (overnight) to give close to quantitative incorporati n. After 35 deprotection of the N-terminal Boc group, a large portion of H-[Taeg]:-Lys(ClZ)-MBHA was taken out for HF cl avage. Boc-

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[Ta g]_m-Lys(ClZ)-MBHA resin was assembled by d ubl in situ DCC c upling of 0.16 M BocTaeg-OH, t g ther with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol" 9). Before coupling of the final residue, a small portion of H-[Taeg],-5 Lys(ClZ)-MBHA was taken out for HF cleavage.

(b) Cleavage, Purification, and Identification f H-[Taeg],-Lys-NH2.

The protected Boc-[Taeg].-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 14.0 mg f 10 crude material upon HF cleavage of 52.4 mg dry H-Ta g].-Lys(ClZ)-MBHA resin. The crude product was not purified (about 99% purity).

- (c) Cleavage, Purification, and Identification of H-[Taeg],-Lys-NH.
- The protected Boc-[Taeg],-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 5.2 mg of crude material upon HF cleavage of 58.4 mg dry H-Taeg],-Lys(ClZ)-MBHA resin.
- (d) Cleavage, Purification, and Identification f H-20 [Taeg],-Lys-NH.

The protected Boc-[Taeg]:-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 114 mg of crude material upon HF cleavage of about 604 mg dry H-Taeg]:-Lys(ClZ)-MBHA resin.

(e) Cleavage, Purification, and Identification f H[Taeg].-Lys-NH.

The protected Boc-[Taeg],-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 19.3 mg f crude material upon HF cleavage of 81.0 mg dry H-Taeg],-30 Lys(ClZ)-MBHA resin.

(f) Cleavage, Purification, and Identification f H-[Taeg],-Lys-NH,.

The protected Boc-[Taeg]_N-Lys(Cl2)-MBHA resin was treated as described in Example 17c to yield about 141 mg of 35 crude mat rial up n HF cl avage f about 417 mg dry H-Ta g]_N-Lys(Cl2)-MBHA resin.

(g) Synthetic Pr t c 1 8 (G neral Pr t c 1)

(1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 3 x 1 min and 1 x 30 min; (2) washing with CH_2Cl_2 , 6 x 1 min; (3) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 3 x 2 min; (4) 5 washing with CH2Cl2, 6 x 1 min, and drain for 1 min; (5) at some stages of the synthesis, 2-5 mg sample of PNA-resin is taken out and dried thoroughly for a ninhydrin analysis to determine the substitution; (6) addition of Boc-protected PNA monomer (Pfp ester); the coupling reaction was allowed to 10 proceed for a total of X hrs shaking; (7) washing with DMF, 1 \times 2 min; (8) washing with CH_2Cl_2 , 4 \times 1 min; (9) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 2 x 2 min; (10) washing with CH₂Cl₂, 6 x 1 min; (11) occasionally, 2-5 mg sample of protected PNA-resin is taken out and dried 15 thoroughly for a ninhydrin analysis to determine the extent of coupling; (12) at some stages of the synthesis, unreact d amino groups are blocked by acetylation with a mixture of acetic anhydride/pyridine/CH₂Cl₂ (1:1:2, v/v/v) for 2 h followed by washing with CH2Cl2, 6 x 1 min, and, occasionally, 20 minhydrin analysis.

EXAMPLE 88

Solid-Phase Synthesis of H-[Taeg]4-Caeg-[Taeg]5-Lys-NH,.

(a) Stepwise Assembly of Boc-[Taeg]4-C[Z]aeg-[Taeg]5-25 Lys(ClZ)-MBHA Resin.

About 1 g of wet Boc-[Taeg]5-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]4-C[Z]aeg-[Taeg]5-Lys(Cl2)-MBHA resin was assembled by in situ DCC coupling of all residues utilizing 0.16 M of BocC[Z]aeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH2Cl2 or 0.16 M BocTaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH2Cl2 ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed about 98% incorporation of C[Z]aeg and cl se to quantitative incorporation of all the Taeg residues.

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(b) Cleavage, Purificati n, and Id ntification f H-[Taeg]4-C[Z]aeg-[Taeg]5-Lys-NH₂.

The protected Boc-[Taeg]4-C[Z]aeg-[Taeg]5-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 5 22.5 mg of crude material upon HF cleavage of 128.2 mg dry H-[Taeg]4-C[Z]aeg-[Taeg]5-Lys(ClZ)-MBHA resin. Crude product (5.8 mg) was purified to give 3.1 mg of H-[Taeg]4-Caeg-[Taeg]5-Lys-NH₂.

- (c) Synthetic Protocol 9 (General Protocol)
- (1) Boc-deprotection with TFA/CH₂Cl₂ (1:1, v/v), 3 x 1 10 min and 1 x 30 min; (2) washing with CH2Cl2, 6 x 1 min; (3) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 3 x 2 min; (4) washing with CH2Cl2, 6 x 1 min, and drain for 1 min; (5) at some stages of the synthesis, 2-5 mg sample of PNA-resin is 15 taken out and dried thoroughly for a ninhydrin analysis t determine the substitution; (6) addition of Boc-protect d PNA · monomer (free acid) in X ml DMF followed by addition of DCC in X ml CH2Cl2; the coupling reaction was allowed to pr ceed for a total of Y hrs shaking; (7) washing with DMF, 1 x 2 min; 20 (8) washing with CH₂Cl₂, 4 x 1 min; (9) neutralization with DIEA/CH₂Cl₂ (1: 19, v/v), 2 x 2 min; (10) washing with CH₂Cl₂, 6 x 1 min; (11) occasionally, 2-5 mg sample of protected PNAresin is taken out and dried thoroughly for a ninhydrin analysis to determine the extent of coupling; (12) at some 25 stages of the synthesis, unreacted amino groups are blocked mixture a acetylation with anhydride/pyridine/CH2Cl2 (1:1:2, v/v/v) for 2 h followed by washing with CH_2Cl_2 , 6 x 1 min, and, occasionally, ninhydrin analysis.

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(NB = COCH3)

EXAMPLE 89 Solid-Phase Synthesis of H-[Taeg]4-(MBaeg)-[Taeg]5-Lys- NH_2 .

(a) Stepwise Assembly of Boc-[Taeg]4-(NBaeg)-[Ta g]5-35 Lys(Cl2)-MBHA Resin.

Ab ut 1 g of wet Boc-[Taeg]5-Lys(ClZ)-MBHA resin was plac d in a 5 ml SPPS r action vessel. Boc-[Taeg]4-(NBaeg)-

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[Taeg]5-Lys(Cl2)-MBHA resin was assembled by in situ DCC coupling utilizing 0.16 M of Boc(NBaeg)-OH together with 0.16 M DCC in 2.0 ml neat CH₂Cl₂ or 0.16 M BocTaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9").

5 Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The NBaeg residue was coupled three times and the Taeg residues were all coupled once. The synthesis was monitored by the ninhydrin reaction which showed >99% total incorporation of NBaeg (about 88% after the first coupling and about 93% after the second coupling) and close to quantitative incorporation of all the Taeg residues.

(b) Cleavage, Purification, and Identification of H[Taeg]4-(NBaeg)-[Taeg]5-Lys-NH2.

The protected Boc-[Taeg]4-(NBaeg)-[Taeg]5-Lys(Cl2)-MBHA

15 resin was treated as described in Example 17c to yield about

33.6 mg of crude material upon HF cleavage of 108.9 mg dry H
[Taeg]4-(NBaeg)-[Taeg]5-Lys(Cl2)-MBHA resin. Crude product

(20.6 mg) was purified to give 4.6 mg of H-[Taeg]4-(NBaeg)
[Taeg]5-Lys-NH2. For (M+H)+, the calculated m/z value was

20 2683.12 and the measured m/z value was 2683.09.

EXAMPLE 90

Solid-Phase Synthesis of H-[Taeg]4-aeg-[Taeg]5-Lys-NH,.

(a) Stepwise Assembly of Boc-[Taeg]4-aeg-[Taeg]5-25 Lys(Cl2)-MBHA Resin.

About 1 g of wet Boc-[Taeg]5-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]4-aeg-[Taeg]5-Lys(Cl2)-MBHA resin was assembled by in situ DCC single coupling of all residues utilizing: (1) 0.16 M of Bocaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ r (2) 0.16 M BocTaeg-OH together with (2) 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitative incorporation of all the residues.

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(b) Cleavage, Purificati n, and Identifi ati n f H-[Taeg]4-aeg-[Ta g]5-Lys-NH2.

The protected Boc-[Taeg]4-aeg-[Taeg]5-Lys(Cl2)-MBHA resin was treated as described in Example 17c to yield about 5 22.2 mg of crude material upon HF cleavage of 126.0 mg dry H-[Taeg]4-aeg-[Taeg]5-Lys(Cl2)-MBHA resin. Crude product (22.2 mg) was purified to give 7.6 mg of H-[Taeg]4-aeg-[Taeg]5-Lys-NH2. For (M+H)+, the calculated m/z value was 2641.11 and the measured m/z value was 2641.16.

10

EXAMPLE 91

Solid-Phase Synthesis of H-[Taeg]4-Gly-[Taeg]5-Lys-NH,.

- (a) Stepwise Assembly of Boc-[Taeg]4-Gly-[Taeg]5-Lys(ClZ)-MBHA Resin.
- About 1 g of wet Boc-[Taeg]5-Lys(ClZ)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]4-Gly-[Taeg]5-Lys(ClZ)-MBHA resin was assembled by in situ DCC single coupling of all residues utilizing: (1) 0.16 M f BocGly-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ r (2) 0.16 M BocTaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reacti n was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitative incorporation of all the 25 residues.
 - (b) Cleavage, Purification, and Identification of H-[Taeg]4-Gly-[Taeg]5-Lys-NH2.

The protected Boc-[Taeg]4-Gly-[Taeg]5-Lys(Cl2)-MBHA resin was treated as described in Example 18c to yield about 30 45.0 mg of crude material upon HF cleavage of 124.1 mg dry H-[Taeg]4-Gly-[Taeg]5-Lys(Cl2)-MBHA resin. Crude product (40.4 mg) was purified to give 8.2 mg of H-[Taeg]4-Gly-[Taeg]5-Lys-NH₂.

Solid-Phas Synthesis of H-[Taeg]4-Gly2-[Ta g]5-Lys-NH2.

- (a) St pwise Assembly of Boc-[Taeg]4-Gly2-[Taeg]5-Lys(ClZ)-MBHA Resin.
- About 1 g of wet Boc-[Taeg]5-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]4-[C[Z]aeg]2-Taeg-C[Z]aeg-Taeg-C[Z]aeg-Lys(Cl2)-MBHA resin was assembled by in situ DCC single coupling of all residues utilizing: (1) 0.16 M of BocGly-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ or (2) 0.16 M BocTaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitativ incorporation of all the residues.
 - (b) Cleavage, Purification, and Identification of H-[Taeg]4-Gly2-[Taeg]5-Lys-NH2.

The protected Boc-[Taeg]4-Gly2-[Taeg]5-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 32.6 mg of crude material upon HF cleavage of 156.6 mg dry H-[Taeg]4-Gly2-[Taeg]5-Lys(ClZ)-MBHA resin. Crude product (30 mg) was purified to give 7.8 mg of H-[Taeg]4-Gly2-[Taeg]5-Lys-NH2. For (M+H)+, the calculated m/z value was 2655.09 and the measured m/z value was 2655.37.

Solid-Phas Synthesis f H-[Taeg]4-[Ca g]2-Taeg-Ca g-Ta g-Caeg-Lys-NH2.

(a) Stepwise Assembly of Boc-[Taeg]4-[C[Z]aeg]2-Taeg5 C[Z]aeg-Taeg-C[Z]aeg-Lys(ClZ)-MBHA Resin.

About 1.5 g of wet Boc-Lys(Cl2)-MBHA (0.28 mmol.Lys/g) resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg]4-[C[Z]aeg]2-Taeg-C[Z]aeg-Taeg-C[Z]aeg-Lys(Cl2)-MBHA resin was assembled by in situ DCC single coupling of all residu s 10 utilizing: (1) 0.16 M of BocC[Z]-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ or (2) 0.16 M BocTaeg-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monit red 15 by the ninhydrin reaction, which showed close to quantitative incorporation of all the residues.

(b) Cleavage, Purification, and Identification of H[Taeg],-[Caeg],-Taeg-Caeg-Taeg-Caeg-Lys-NH,.

The protected Boc-[Taeg]4-[C[Z]aeg]2-Taeg-C[Z]aeg-Taeg-20 C[Z]aeg-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 52.1 mg of crude material upon HF cleavage of 216.7 mg dry H-[Taeg]4-[C[Z]aeg]2-Taeg-C[Z]a g-Taeg-C[Z]aeg-Lys(ClZ)-MBHA resin. Crude product (30.6 mg) was purified to give 6.2 mg of H-[Taeg]4-[Caeg]2-Taeg-Caeg-25 Taeg-Caeg-Lys-NH2. For (M+H)+ the calculated m/z value was 2747.15 and the measured m/z value was 2746.78.

EXAMPLE 94

Solid-Phase Synthesis of H-Caeg-Taeg-Caeg-Taeg-[Caeg]3-Taeg-30 Caeg-Taeg-Lys-NH2.

(a) Stepwise Assembly of Boc-C[Z]aeg-Taeg-C[Z]aeg-Taeg-[C[Z]aeg-Taeg-C[Z]aeg-Taeg-Lys(ClZ)-MBHA Resin.

About 1.5 g of wet Boc-Lys(ClZ)-MBHA (0.28 mmol Lys/g) resin was placed in a 5 ml SPPS reaction vessel. Boc-C[Z]a g-35 Taeg-C[Z]aeg-Taeg-[C[Z]aeg]3-Taeg-C[Z]aeg-Taeg-Lys(ClZ)-MBHA r sin was assembled by in situ DCC single c upling f all residues utilizing: (1) 0.16 M of BocC[Z]-OH togeth r with

0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ or (2) 0.16 M BocTa g-OH together with 0.16 M DCC in 2.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was 5 monitored by the ninhydrin reaction, which showed close to quantitative incorporation of all the residues.

(b) Cleavage, Purification, and Identification of E-Caeg-Taeg-Caeg-Taeg-Caeg-Taeg-Lys-NH 2.

The protected Boc-C[Z]aeg-Taeg-C[Z]aeg-Taeg-[C[Z]aeg]3
10 Taeg-C[Z]aeg-TaegLys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 56.1 mg of crude material upon HF cleavage of 255.0 mg dry H-C[Z]aeg-Taeg-C[Z]aeg-Taeg-[C[Z]aeg]3-Taeg-C[Z]aeg -TaegLys(ClZ)-MBHA resin. Crude product (85.8 mg) was purified to give 46.2 mg of H-Caeg-Taeg
15 Caeg-Taeg-[Caeg]3-Taeg-Caeg-Taeg-LysNH2. For (M+H)+2 the calculated m/z value was 2717.15 and the measured m/z value was 2716.93.

EXAMPLE 95

- 20 Solid-Phase Synthesis of H-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH2, H-Caeg-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH2, and H-Tyr-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH2.
- (a) Stepwise Assembly of Boc-[Taeg]2-[C(Z)aeg]3[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA Resin, Boc-Caeg-[Taeg]225 [C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA Resin, and B cTyr(BrZ)-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA
 Resin.

About 3 g of wet Boc-Lys(Cl2)-MBHA (0.28 mmol Lys/g) resin was placed in a 20 ml SPPS reaction vessel. B c-30 [Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(Cl2)-MBHA resin was assembled by in situ DCC single coupling of all residues utilizing: (1) 0.16 M of BocC[Z]-OH together with 0.16 M DCC in 3.0 ml 50% DMF/CH2Cl2 or (2) 0.16 M BocTaeg-OH together with 0.16 M DCC in 3.0 ml 50% DMF/CH2Cl2 ("Synthetic Protoc l 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitative

incorporation of all the residu s. After deprotecti n of the N-terminal Boc group, half of the PNA-resin was coupled quantitatively onto Tyr(BrZ)-OH and a small portion was coupled quantitatively onto one more Caeg residue. Both 5 couplings employed the above-mentioned synthetic protocol.

(b) Cleavage, Purification, and Identification of H-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH2.

The protected Boc-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 50.9 mg of crude material upon HF cleavage of 182.5 mg dry H-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA resin. Crude product (50.9) mg was purified to give 13.7 mg of H-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-LysNH2. For (M+H)+ the calculated m/z value was 2466.04; the m/z value was not measured.

(c) Cleavage, Purification, and Identification of H-Tyr-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH2.

The protected Boc-Tyr(BrZ)-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA resin was treated as described in 20 Example 17c to yield about 60.8 mg of crude material upon HF cleavage of 188.8 mg dry H-Tyr(BrZ)-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA resin. Crude product (60.8 mg) was purified to give 20.7 mg of H-Tyr-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-LysNH2. For (M+H)+ the calculated m/z value 25 was 2629.11 and the measured m/z value was 2629.11.

(d) Cleavage, Purification, and Identification of H-Caeg-[Taeg]2-[Caeg]3-[Taeg]2-[Caeg]2-Lys-NH₂.

The protected Boc-C(Z)aeg-[Taeg]2-[C(Z)aeg]3-[Taeg]2-30 [C(Z)aeg]2-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 11.7 mg of crude material upon HF cleavage of 42.0 mg dry H-C(Z)aeg-[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(ClZ)-MBHA resin. Crude product (11.6 mg) was purified to give 3.1 mg of H-Caeg-[Taeg]2-[Caeg]3-[Taeg]2-35 [Caeg]2-LysNH2. For (M+H)+ the calculated m/z value was 2717.15; th m/z value was n t measured.

Solid-Phase Synthesis of H-[Caeg]2-[Taeg]2-[Caeg]3-[Taeg]2-Lys-NH2,

H-Taeg-[Caeg]2-[Taeg]2-[Caeg]3-[Taeg]2-Lys-NH₂, and H-Tyr-5 [Caeg]2-[Taeg]2-[Caeg]3-[Taeg]2-Lys-NH₂.

(a) Stepwise Assembly of Boc-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys(ClZ)-MBHA Resin, Boc-Taeg-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys(ClZ)-MBHA Resin, and Boc-Tyr(BrZ)-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys(ClZ)-MBHA Resin.

About 3 g of wet Boc-Lys(Cl2)-MBHA (0.28 mmol Lys/g) resin was placed in a 20 ml SPPS reaction vessel. B c[Taeg]2-[C(Z)aeg]3-[Taeg]2-[C(Z)aeg]2-Lys(Cl2)-MBHA resin was assembled by in situ DCC single coupling of all residues

15 utilizing: (1) 0.16 M of BocC[Z]-OH together with 0.16 M.DCC in 3.0 ml 50% DMF/CH₂Cl₂ or (2) 0.16 M BocTaeg-OH together with 0.16 M DCC in 3.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed for a total of 20-24 hrs with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitative incorporation of all the residues. After deprotection of the N-terminal Boc group, half of the PNA-resin was coupled quantitatively onto Tyr(BrZ)-OH and a small portion was coupled quantitatively onto one more Taeg residue. Both 25 couplings employed the above-mentioned synthetic protocol.

(b) Cleavage, Purification, and Identification of H-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys-NH₂.

The protected Boc-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3[Taeg]2-Lys(ClZ)-MBHA resin was treated as described in

30 Example 17c to yield about 57.6 mg of crude material upon HF
cleavage of 172.7 mg dry H-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3[Taeg]2-Lys(ClZ)-MBHA resin. Crude product (57.6 mg) was
purified to give 26.3 mg of H-[Caeg]2-[Taeg]2-[Caeg]3-[Taeg]2Lys-NH2. For (M+H)+ the calculated m/z value was 2466.04; the

35 m/z value was not measured.

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(c) Cl avage, Purificati n, and Identificati n f H-Tyr-[C(2)aeg]2-[Taeg]2-[C(2)aeg]3-[Taeg]2-Lys-NH, .

The protected B c-Tyr(BrZ)-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 57.6 mg of crude material upon HF cleavage of 172.7 mg dry H-Tyr(BrZ)-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys(ClZ)-MBHA resin. Crude product (47.1 mg) was purified to give 13.4 mg of H-Tyr-[Caeg]2-[Taeg]2-[Caeg]3-[Taeg]2-Lys-NH2. For (M+H)+ the calculated m/z value was 2629.11 and the measured m/z value was 2629.11.

(d) Cleavage, Purification, and Identification of H-Taeg-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3-[Taeg]2-Lys-NH₂.

The protected Boc-Taeg-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]315 [Taeg]2-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 53.4 mg of crude material upon HF cleavage of 42.4 mg dry H-Taeg-[C(Z)aeg]2-[Taeg]2-[C(Z)aeg]3[Taeg]2-Lys(Cl Z)-MBHA resin. Crude product (11.9 mg) was purified to give 4.3 mg of H-Taeg-[Caeg]2-[Taeg]2-[Caeg]320 [Taeg]2-Lys-NH2. For (M+H)+ the calculated m/z value was 2732.15; the m/z value was not measured.

(c) Synthetic Protocol 10 (General Protocol)

Same protocol as "Synthetic Protocol 9", except that DCC has been replaced with DIC.

25

EXAMPLE 97

SYNTHESIS OF THE BACKBONE MOIETY FOR SCALE UP BY REDUCTIVE AMINATION

(a) Preparation of (bocamino) acetaldehyde.

3-Amino-1,2-propanediol(80.0 g; 0.88 mol) was dissolved in water (1500 ml) and the solution was cooled to 4°C, whereafter Boc anhydride (230 g; 1.05 mol) was added at once. The solution was gently heated to room temperature with a water bath. The pH was kept at 10.5 by the dropwise addition of sodium hydroxide. Over the course of the reaction a total f 70.2 g NaOH, dissolv d in 480 ml water, was added. After stirring vernight, ethyl acetat (1000 ml) was added and the

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mixture was cooled to 0°C and the pH was adjusted to 2.5 by the addition f 4 M hydrochloric acid. The thyl acetate layer was removed and the acidic aqueous solution was extracted with more ethyl acetate (8x500 ml). The combined 5 ethyl acetate solution was reduced to a volume of 1500 ml using a rotary evaporator. The resulting solution was washed with half saturated potassium hydrogen sulphate (1500 ml) and then with saturated sodium chloride. It then was dried over magnesium sulphate and evaporated to dryness, in vacuo. 10 Yield. 145.3 g (86%)

3-Bocamino-1,2-propanediol (144.7 g; 0.757 mol) was suspended in water (750 ml) and potassium periodate (191.5 g; 0.833 mol) was added. The mixture was stirred under nitr gen for 2.5 h and the precipitated potassium iodate was rem ved 15 by filtration and washed once with water (100 ml). Th aqueous phase was extracted with chloroform (6x400 ml). chloroform extracts were dried and evaporated to dryness, in Yield 102 g (931) of an oil. . vacuo. (bocamino) acetaldehyde was purified by kugelrohr distillati n 20 at 84°C and 0.3 mmHg in two portions. The yield 79 g (77%) of a colorless oil.

(b) Preparation of (N'-bocaminoethyl)glycine methyl ester

Palladium on carbon (10%; 2.00 g) was added to a solution of (bocamino) acetaldehyde (10.0 g; 68.9 mmol) in methanol (150 ml) at 0°C. Sodium acetate (11.3 g; 138 mmol) in methanol (150 ml), and glycine methyl ester hydrochloride (8.65 g; 68.9 mmol) in methanol (75 ml) then were added. The mixture was hydrogenated at atmospheric pressure for 2.5 h, 30 then filtered through celite and evaporated to dryness, in vacuo. The material was redissolved in water (150 ml) and the pH was adjusted to 8.0 with 0.5 N NaOH. The aqueous solution was extracted with methylene chloride (5 x 150 ml). The combined extracts were dried over sodium sulphate and evap rated to dryness, in vacuo. This resulted in 14.1 g (88%) f (N'-bocaminoethyl)glycine methyl ester. The crude material was purified by kugelrohr destination at 120°C and

0.5 mmHg to give 11.3 g (70%) of a colorless oil. The product had a purity that was higher than the material produced in example 26 according to tlc-analysis (10% methanol in methylene chloride).

Alternatively, sodium cyanoborohydride can be used as reducing agent instead of hydrogen (with Pd(C) as catalyst), although the yield (42%) was lower.

- (c) Preparation of (N'-bocaminoethyl)glycine ethyl ester.
- The title compound was prepared by the above procedure with glycine ethyl ester hydrochloride substituted for glycine methyl ester hydrochloride. Also, the solvent used was ethanol. The yield was 78%.

15 EXAMPLE 98

Solid-Phase Synthesis of H-Tyr-[Taeg] n-Lys-NH,

(a) Stepwise Assembly of Boc-Tyr(BrZ)-[Taeq] $_{\rm H}$ -Lys(ClZ)-MBHA Resin.

About 0.2 g of wet Boc-[Taeg]_W-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-Tyr(BrZ)-[Taeg]_W-Lys(Cl2)-MBHA resin was assembled by standard in situ DCC coupling utilizing 0.32 M of BocCTyr(BrZ)-OH together with 0.32 M DCC in 3.0 ml neat CH₂Cl₂ overnight. The ninhydrin reaction showed about 97% incorporation of BocTyr(BrZ).

25 (b) Cleavage, Purification, and Identification of H-Tyr-[Taeg]_w-Lys-NH₂.

The protected Boc-Tyr(BrZ)-[Taeg]_N-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 5.5 mg of crude material upon HF cleavage of 20.7 mg dry H-Tyr(BrZ)-30 [Taeg]_N-Lys(ClZ)-MBHA resin. The crude product was purified to give 2.5 mg of H-Tyr-[Taeg]_N-Lys-NH₂.

Solid-Phase Synth sis of Dansyl-[Taeg],-Lys-NH,

- (a) Stepwise Assembly of Dansyl-[Taeg],-Lys(ClZ)-MBHA Resin.
- About 0.3 g of wet Boc-[Taeg]_N-Lys(ClZ)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Dansyl-[Taeg]_N-Lys(ClZ)-MBHA resin was assembled by coupling of 0.5 M dansyl-Cl in 2.0 ml neat pyridine overnight. The ninhydrin reaction showed about 95% incorporation of dansyl.
- 10 (b) Cleavage, Purification, and Identification of Dansyl-[Taeg],-Lys-NH2.

The protected dansyl-[Taeg]₁₀-Lys(Cl2)-MBHA resin was treated as described in Example 17c to yield about 12 mg of crude material upon HF cleavage of 71.3 mg dry dansyl-[Taeg]₁₀-15 Lys(Cl2)-MBHA resin. The crude product was purified to give 5.4 mg of dansyl-[Taeg]₁₀-Lys-NH₂.

EXAMPLE 100

Solid-Phase Synthesis of Gly-Gly-His-[Taeg],-Lys-NH,

20 (a) Stepwise Assembly of Boc-Gly-Gly-His(Tos)-[Taeg],-Lys(ClZ)-MBHA Resin.

About 0.05 g of Boc-[Taeg]₁₀-Lys(Cl2)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-Gly-Gly-His(Tos)-[Taeg]₁₀-Lys(Cl2)-MBHA resin was assembled by standard double 25 in situ DCC coupling of Boc-protected amino acid (0.1 M) in 2.5 ml 25% DMF/CH₂Cl₂, except for the first coupling f BocHis(Tos), which was done by using a preformed symmetrical anhydride (0.1M) in 25% DMF/CH₂Cl₂. All couplings were performed overnight and ninhydrin reactions were not carried out.

(b) Cleavage, Purification, and Identification of Gly-Gly-His-[Taeg],-Lys-NE₂.

The protected Boc-Gly-Gly-His(Tos)-[Taeg]₁₀-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield 35 about 10.3 mg of crude material (about 40% purity) upon HF cleavage of 34.5 mg dry Boc-Gly-Gly-His(Tos)-[Taeg]₁₀-Lys(ClZ)-

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MBHA resin. A small p rti n f the crude product (taken ut bef re ly philizati n) was purified t giv 0.1 mg f Gly-Gly-His-[Ta g],-Lys-NH,.

5 EXAMPLE 101

Solid-Phase Synthesis of H-[Taeg],-[Caeg],-NH,.

(a) Stepwise Assembly of Boc-[Taeg],-[C(2)aeg],-MBHA Resin.

About 0.2 g of MBHA resin was placed in a 3 ml SPPS 10 reaction vessel and neutralized. The loading was determined to be about 0.64 mmol/g. Bocc(Z)aeg-OPfp was coupled onto the resin using a concentration of 0.13 M in 2.5 ml 25% phenol//CH₂Cl₂. The ninhydrin analysis showed a coupling yield of about 40%. The remaining free amino groups were acetylated 15 as usual. Boc-[Taeg];-[C(Z)aeg];-MBHA resin was assembled by single in situ DCC coupling of the next residue utilizing 0.11 M of Bocc(Z)aeg-OH together with 0.11 M DCC in 2.5 ml 50% DMF/CH;Cl; and by coupling with 0.13 M BocTaeg-OPfp in n at CH.Cl. for the remaining residues ("Synthetic Protocol 8"). 20 Each coupling reaction was allowed to proceed with shaking The synthesis was monitored by the ninhydrin

reaction, which showed close to quantitative incorporation of all the residues.

(b) Cleavage, Purification, and Identification f H-25 [Taeg],-[Caeg],-NH2.

The protected Boc-[Taeg],-[C(Z)aeg],-MBHA resin was treated as described in Example 17c to yield about 21.7 mg of crude material (>80% purity) upon HF cleavage of 94.8 mg dry H-[Taeq]:-[C(Z)aeq]:-MBHA resin. Crude product (7.4 mg) was 30 purified to give 2.0 mg of H-[Taeg],-[Caeg],-NH, (>99% purity).

EXAMPLE 102

Solid-Phase Synthesis of H-[Taeg],-Caeg-[Taeg],-NH,.

(a) Stepwise Assembly of Boc-[Taeg],-C(Z)aeg-[Ta g],-35 MBHA Resin.

f the above-mentioned MBHA resin was About 0.2 g plac d in a 5 ml SPPS reacti n vess l and neutralized. Boc[Taeg],-C(Z)aeg-[Taeg].-MBHA resin was assembled by single in situ DCC coupling of th C(Z)aeg residue utilizing 0.13 M f BocC[Z]aeg-OH together with 0.13 M DCC in 2.5 ml 50% DMF/CH,Cl, and by coupling the Taeg residues with 0.13 M BocTaeg-OPfp in 2.5 ml neat CH₂Cl₂. Each coupling reaction was allowed to proceed with shaking overnight. The synthesis was monit red by the ninhydrin reaction, which showed close to quantitative incorporation of all the residues.

(b) Cleavage, Purification, and Identification of H-10 [Taeg],-Caeg-[Taeg],-NH2.

The protected Boc-[Taeg],-C(Z)aeg-[Taeg],-MBHA resin was treated as described in Example 17c to yield about 44.4 mg of crude material upon HF cleavage of about 123 mg dry H-[Taeg],-C(Z)aeg-[Taeg],-MBHA resin. Crude product (11.0 mg) was purified to give 3.6 mg of H-[Taeg],-Caeg-[Taeg],-NH,.

EXAMPLE 103

Solid-Phase Synthesis of H-Taeg-Caeg-[Taeg],-LysNH, .

(a) Stepwise Assembly of Boc-Taeg-C(2)aeg-[Taeg],-20 Lys(Cl2)-MBHA Resin.

About 0.3 g of wet Boc-[Taeg];-Lys(Cl2)-MBHA resin was placed in a 3 ml SPPS reaction vessel. Boc-Taeg-C(Z)a g-[Taeg];-Lys(Cl2)-MBHA resin was assembled by single in situ DCC coupling overnight of the C(Z)aeg residue ("Synthetic Protocol" 9) utilizing 0.2 M of BocC[Z]aeg-OH together with 0.2 M DCC in 2.5 ml 50% DMF/CH;Cl; (incorporation was about 80% as judged by ninhydrin analysis; remaining free amino groups were acetylated) and by overnight coupling the Taeg residue with 0.15 M BocTaeg-OPfp in 2.5 ml neat CH;Cl; (nearly 30 quantitatively).

(b) Cleavage, Purification, and Identification of H-Taeg-Caeg-[Taeg],-LysNH1.

The protected Boc-Taeg-C(Z)aeg-[Taeg].-Lys(C1Z)-MBHA resin was treated as described in Example 17c to yield about 35 22.3 mg f crude material up n HF cl avag of about 76.5 mg dry H-Taeg-C(Z)aeg-[Taeg].-Lys(C1Z)-MBHA resin. Crude product

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(6.7 mg) was purified t giv 2.6 mg f H-Taeg-Caeg-[Ta g].-LysNH₂. Fr (M+H) th calculated m/z value was 2792.15 and the measured m/z value was 2792.21.

5 EXAMPLE 104

Caeg-[Taeg],-Lys-NH,.

Solid-Phase Synthesis of H-Caeg-[Taeg],-Lys-NH, and H-[Ta g],-Caeg-[Taeg],-Lys-NH,.

- (a) Stepwise Assembly of Boc-[Taeg]₁-C(2)aeg-[Ta g]_i-Lys(Cl2)-MBHA Resin.
- About 0.5 g of wet Boc-[Taeg];-Lys(ClZ)-MBHA resin was placed in a 5 ml SPPS reaction vessel. Boc-[Taeg];-C(Z)a g-[Taeg];-Lys(ClZ)-MBHA resin was assembled by single in situ DCC coupling of all residues utilizing: (1) 0.12 M f BocC[Z]aeg-OH together with 0.12 M DCC in 3.0 ml 50% DMF/CH;Cl;
- or (2) 0.12 M BocTaeg-OH together with 0.12 M DCC in 3.0 ml 50% DMF/CH₂Cl₂ ("Synthetic Protocol 9"). Each coupling reaction was allowed to proceed overnight with shaking. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitative incorporation of all th 20 residues. During the synthesis, a small portion of H-C(Z)aeg-
 - [Taeg],-Lys(Cl2)-MBHA resin was taken out for HF cleavage.

 (b) Cleavage, Purification, and Identification f H-
- The protected Boc-C[Z]aeg-[Taeg];-Lys(ClZ)-MBHA resin 25 was treated as described in Example 17c to yield about 3.0 mg of crude material upon HF cleavage of 37.5 mg dry H-C[Z]aeg-[Taeg];-Lys(ClZ)-MBHA resin. About 0.7 mg of the crude product was purified to give about 0.5 mg of H-Caeg-[Taeg];-Lys-NH;.

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(c) Cleavage, Purification, and Identification f H-[Taeg];-Caeg-[Taeg];-Lys-NH;.

The protected Boc-[Taeg];-C[Z]aeg-[Taeg];-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 35 37.7 mg of crude material upon HF cleavage f 118.6 mg dry H[Taeg];-C[Z]aeg-[Ta g];-Lys(ClZ)-MBHA r sin.

Solid-Phase Synth sis f H-[Ca g],-Lys-NH2, H-[Ca g],-Lys-NH2, H-[Caeg],-Lys-NH2, and H-[Caeg],-Lys-NH2

(a) Stepwise Assembly of Boc-[C(2)aeg] $_{\rm H}$ -Lys(C12)-MBHA 5 Resin and Shorter Fragments.

About 5 g of wet Boc-Lys(Cl2)-MBHA resin (substitution = 0.3 mmol Lys/g) was placed in a 30 ml SPPS reaction vessel. Boc-[C(Z)aeg]_n-Lys(ClZ)-MBHA resin was assembled by single in situ DCC coupling of the first three residues with 0.1 M of 10 Bocc(Z)aeg-OH together with 0.1 M DCC in 10 ml 50% DMF/CH,Cl, ("Synthetic Protocol 9") and by single in situ DIC coupling of the remaining seven residues with 0.1 M of BocC(Z)aeg-OH together with 0.1 M DIC in 10 ml 50% DMF/CH,Cl, ("Synthetic Protocol 10"). All the coupling reactions were allowed to 15 proceed overnight. The synthesis was monitored by the ninhydrin reaction, which showed close to quantitativ incorporation of all residues. During the synthesis, portions of the shorter fragments H-[C(Z)aeg],-Lys(ClZ)-MBHA resin, H- $[C(Z) aeg]_{\bullet}-Lys(ClZ)-MBHA$ resin, $H-[C(Z) aeg]_{\bullet}-Lys(ClZ)-MBHA$ 20 resin, $H-[C(Z)aeg]_{\bullet}-Lys(ClZ)-MBHA$ resin, and $H-[C(Z)aeg]_{\bullet}-$ Lys(ClZ)-MBHA resin were taken out for HF cleavage.

(b) Cleavage, Purification, and Identification of H-[Caeg],-Lys-NH.

The protected Boc-[C(E)aeg],-Lys(ClZ)-MBHA resin was treated
25 as described in Example 17c to yield about 10.8 mg of crude
material upon HF cleavage of 60.1 mg dry H-[C(Z)aeg],Lys(ClZ)-MBHA resin.

- (c) Cleavage, Purification, and Identification of H-[Caeg],-Lys-NH,.
- The protected Boc-[C(Z)aeg].-Lys(C1Z)-MBHA resin was treated as described in Example 17c to yield about 13.4 mg of crude material upon HF cleavage of 56.2 mg dry H-[C(Z)aeg].-Lys(C1Z)-MBHA resin.

(d) Cleavage, Purifi ati n, and Identificati n f H-[Caeg],-Lys-NH,.

The protected Boc-[C(2)aeg]:-Lys(Cl2)-MBHA resin was treat d as described in Example 17c to yield about 16.8 mg of crud 5 material upon HF cleavage of 65.6 mg dry H-[C(2)aeg]:-Lys(Cl2)-MBHA resin.

(e) Cleavage, Purification, and Identification of H-[Caeg]_u-Lys-NH₂.

The protected Boc-{C(Z)aeg}_N-Lys(ClZ)-MBHA resin was 10 treated as described in Example 17c to yield about 142.4 mg of crude material upon HF cleavage of 441 mg dry H-{C(Z)aeg}_N-Lys(ClZ)-MBHA resin.

EXAMPLE 106

- 15 Solid-Phase Synthesis of H-[Taeg];-Caeg-[Taeg];-Caeg-[Taeg],-Lys-NH,
 - (a) Stepwise Assembly of Boc-[Taeg]₂-C(Z)aeg-[Ta g]₂-C(Z)aeg-[Taeg]₄-Lys(ClZ)-MBHA Resin.

About 0.3 g of wet H-[Taeg],-C(Z)aeg-[Taeg],-Lys(ClZ)-20 MBHA resin from the earlier synthesis of Boc-[Taeg],-C(Z)a g-[Taeg].-Lys(ClZ)-MBHA resin was placed in a 5 ml SPPS reacti n After coupling of the next residue five times, a total incorporation of BocC(Z)aeg of 87% was obtained. The five repeated couplings were carried out with 0.18 M 25 BocC(Z)aeg-OPfp in 2 ml of TFE/CH₂Cl₂ (1:2, V/V), 2 ml f TFE/CH,Cl, (1:2, v/v), 2 ml of TFE/CH,Cl, (1:2, v/v) with two drops of dioxane and two drops of DIEA (this condition gave only a few per cent coupling yield), 2 ml of TFE/CH2Cl2 (1:2, v/v) plus 0.5 g phenol, and 1 ml of CH₂Cl₂ plus 0.4 g f The two final Taeg residues were 30 phenol, respectively. incorporated close to quantitatively by double couplings with 0.25 M BocTaeg-OPfp in 25% phenol/CH2Cl2. All couplings were allowed to proceed overnight.

(b) Cl awage, Purification, and Identification f H[Taeg];-Ca g-[Ta g];-Caeg-[Ta g],-Lys-NH,

The protected Boc-[Taeg];-C(Z)aeg-[Taeg];-C(Z)aeg-[Taeg],-Lys(ClZ)-MBHA resin was treated as described in Example 17c to yield about 7 mg of crude material upon HF cleavage of 80.7 mg dry H-[Taeg];-C(Z)aeg-[Taeg];-C(Z)aeg-[Taeg],-Lys(ClZ)-MBHA resin. The crude product was purified to give 1.2 mg of H-[Taeg];-Caeg-[Taeg];-Caeg-[Taeg],-Lys-NH; (>99.9% purity).

10 EXAMPLE 107

SYNTHESIS OF A PNA WITH TWO ANTI PARALLEL STRANDS TIED TOGETHER

Synthesis of H-[Taeg]-[Taeg]-[Gaeg]-[Taeg]-[Taeg][Taeg]-[6-AHA]-[aeg]-[6-AHA]-[Taeg]-[Taeg]-[Taeg]-[Aaeg]-[
Taeg]-[Taeg]-[Taeg]-LYS-NH₂. (6-AHA = 6-aminohexanoic acid)
(Pigure 26)

The protected PNA was assembled onto a Boc-Lys(Cl2) modified MBHA resin with a substitution of approximately 0.30 mmol/g. Capping of uncoupled amino groups was only carried out before the incorporation of the BocGaeg-OH monomer. Synthesis was initiated on 1.00 g (dry weight) of preswollen (overnight in DCM) and neutralized Boc-Lys(Cl2)-MBHA resin. The incorporation of the monomers followed the protocol of Example 32 and Example 71. The coupling reaction was monitored by qualitative ninhydrin reaction (kaiser test). In case of a positive Kaiser test, the coupling reaction was repeated until the test showed no coloration of the beads. Final deprotection, cleavage from support, and purification were performed according to standard procedures.

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EXAMPLE 108

Alternative protecting group strategy for PNA-synthesis (Figure 27).

- (a) Synthesis of test compounds.
- 2-amino-6-O-benzyl purine. To a solution of 2.5 g (0.109 mol) of sodium in 100 ml f benzyl alcoh 1 was added 10.75 g (0.063 mol) f 2-amino-6-chloropurine. The mixture

was stirr d f r 12 h at 120 0°C. The s lution was c oled to r om temperature and neutralized with acetic acid and extracted with 10 portions of 50 ml of 0.2 N sodium hydroxide. The collected sodium hydroxide phases were washed with 100 ml 5 of diethyl ether and neutralized with acetic acid, wher by precipitation starts. The solution was cooled to 0°C and the filtrati n. precipitate VAS collected by yellow Recrystallization from ethanol gave 14.2 g 92% of pure whit crystals of the target compound. 1H-NMR (250 MHz--DMSO-d6) 10 d ppm: 8-H, 7.92; benzyl aromatic, 7.60-7.40; 2NH,, 6.36; benzyl CH2, 5.57.

(2-amino-6-O-benzyl purinyl) methylethanoate. A mixture of 5 g (0.0207 mol) of 2-amino-6-O-benzyl-purine, 30 ml of DMF and 2.9 g (0.021 mol) of potassium carbonate was stirred at room temperature. Methyl bromoacetate (3.2 g; 1.9 ml; 0,0209 mol) was added dropwise. The solution was filtrated after 4 h and the solvent was removed under reduced pressure (4 mmHg, 40°C). The residue was recrystallized two times from thyl acetate to give 3.7 g (57%) of the target compound. 1H-NMR (250 MHz, DMSO-d6) d ppm: 8-H, 7.93; benzyl aromatic 7.4-7.6; 2-NH₂, 6.61; benzyl CH2, 5.03; CH2, 5.59; OCH3, 3.78.

ethanoate. To a solution of 0.5 g (1.6 mmol) of (2-amino-6-0-benzyl purinyl) methyl ethanoate in 25 ml methylene chl rid

25 was added 0.53 g (1.62 mmol) of p-toluenesulfonic anhydride
and 0.22 g (1.62 mmol) of potassium carbonate. The mixtur
was stirred at room temperature. The mixture was filtered and
the solvent was removed at reduced pressure (15 mmHg, 40°C).
Diethyl ether was added to the oily residue. The resulting
30 solution was stirred overnight, whereby the target compound
(0.415 mg; 55%) precipitated and was collected by filtrati n.
1H-NMR (250 MHz, DMSO-d6) d ppm: 8-H, 8.97; aromatic 7.2-7.8;
benzyl CH2, 5,01; CH2, 4.24; OCH3, 3.73; CH3, 2.43.

(b) Stability of the tosyl protected base-residue in 35 TFA and HF.

Th material was subj ct d to the standard depr t cti n conditions (TFA-depr tecti n) and th final cleavage

conditions with HF. The products were then subjected to HPLC-analysis using a 4 μ RCM 8x10 Nova pack—lumn and solvents A (0.1% TFA in water) and B (0.1% TFA in acetonitrile) according to the following time gradient with a flow f 2 ml/min.

	Time	* A	₹B
	0	100	Ö
	5	100	Ŏ
	35	0	100
10	37	0	100
	39	100	0

The following retention times were found: (a) Compound 1: 30.77 min; (b) compound 2: 24.22 min; and (c) compound 3: 11.75 min. The analysis showed that the O6-benzyl group was removed both by TFA and HF, whereas there was no cleavage f the tosyl group in TFA, but quantitative removal in HF under the standard cleavage conditions.

EXAMPLE 109

20. 5-Bromouracil-N1-methyl acetate

5-Bromouracil (5.00 g; 26.2 mmol) and potassium carbonate (7.23 g; 52.3 mmol) were suspended in DMF (75 ml). Methyl bromoacetate (2.48 ml; 26.1 mmol) was added over a period of 5 min. The suspension was stirred for 2 h at r om 25 temperature, and then filtered. The solid residue was washed twice with DMF, and the combined filtrates were evaporated to dryness, in vacuo. The residue was an oil containing the title compound, DMF and some unidentified impurities. It is not necessary to purify the title compound before hydrolysis. 30 H-NMR (DMSO-d4, 250 MHz); 8.55 (impurity); 8.27 (CBr=CHN); 8.02 (impurity); 4.76 (impurity); 4.70 (impurity); 4.62 (NCH_2COOCH_3) ; 3.78 $(COOCH_3)$; 2.96 (DMF); 2.80 (DMF). ¹⁵C-NMR (DMSO-d₆, 250 MHz); 168.8 (COOCH₃); 172.5 (CH=CBrCON); 161.6 (DMF); 151.9 (NCON); 145.0 (CO-CBr=CHN); 95.6 (COCBr=CHN); (impurity); 52.5 (OCH3); 49.7 (impurity); (NCH,COOMe); 43.0 (impurity); 36.0 (DMF). UV(Methanol; __nm); 226; 278. IR (KBr;cm⁻¹_; 3158s (_NH); 1743vs (_C=0, COOMe); 1701vs (_C=0, CONH); 1438vs (d CH, CH₃O); 1223vs (_ C-O,

COOMe); 864 m (∂ CH, Br=C-H). FAB-MS m/z (assignment): 265/263 (M+H).

EXAMPLE 110

5 (5-Bromouracil) acetic acid

Water (30 ml) was added to the oil of the crude product from Example 109 and the mixture was dissolved by adding sodium hydroxide (2M, 60 ml). After stirring at 0°C for 10 min, hydrochloric acid (4M, 45 ml) was added to pH=2 10 and the title compound precipitated. After 50 min, the s lid residue was isolated by filtration, washed once with c ld water, and then dried in vacuo over sicapent. Yield: 2.46 g (38%). Mp, 250°-251°C. Anal. for CaHaBrN2O4. Found (calc.): C: 28.78 (28.94); H: 2.00 (2.02); Br: 32.18 (32.09); N: 11.29 15 (11.25). H-NMR (DMS.-d, 250 MHz): 12,55 (1H.s,COOH); 11.97 (1H,s,NH); 8.30 (1H,s,C=C-H); 4.49 (2H,s,NCH2COOH). 13 C-NMR (DMSO-d₆, 250 MHz); 169.4 (COOH); 159.8 (NHCOCBr=CH); 150.04 (NCON); 145.8 (COCBr=CHN); 94.6 (COCBr=CHN); 48.8 (NCH2COOH). UV (Methanol; _mxnm); 226; 278. IR (KBr; cm⁻¹); 3187s (_NH); 20 1708vs (_C=0,COOH); 1687vs; 1654VS (_C=0, CONH); 1192s (_C=0, COOH); 842 m (8 CH, Br-C=C-H). FAB-MS m/z (assignment, relative intensity); 251/249 (M + H,5).

EXAMPLE 111

25 N-(Boc-aminoethyl)-N-(5-bromouracil)methylenecarbonoylglycine ethyl ester

Boc-aminoethylglycine ethyl ester (1.80 g; 7.30 mmol) was dissolved in DMF (10 ml). Dhbt-OH (1.31 g; 8.03 mmol) was added, whereby a precipitate was formed. DMF (2 x 10 ml) was added until the precipitate was dissolved. The product f Example 110 (2.00 g; 8.03 mmol) was added slowly to avoid precipitation. Methylene chloride (30 ml) was added, and the mixture was cooled to 0°C and then filtered. The precipitat, DCU, was washed twice with methylene chloride. To the mixture was washed with half saturat d NaHCO3-s luti n (3 x 100 ml, H2O:saturated NaHCO3-s lution 1:1 v/v), then with

dilut KHSO,-solution (2 x 100 ml, H,O:saturated KHSO,-solution 4:1 V/V, and finally with saturated NaCl-solution (1 x 100 ml). The organic phase was dried over magnesium sulphate, filtered, and evaporated to dryness in vacuo (about 15 mmHg 5 and then about 1 mmHg). The residue was suspended in methylene chloride (35 ml), stirred for 45 min at room temperature, and filtered (the precipitate was DCU). Petroleum ether (2 volumes) was added dropwise to the filtrate at 0°C, whereby an oil precipitated. The liquor was decanted 10 and the remaining oil dissolved in methylene chloride (20-50 ml). Precipitated was effected by the addition of petroleum ether (2 volumes). This procedure was repeated 5 times until an impurity was removed. The impurity can be seen at TLC with 10% MeOH/CH,Cl, as the developing solvent. The resulting il 15 was dissolved in methylene chloride (25 ml) and evaporated to dryness in vacuo, which caused solidification of the title compound. Yield: 2.03 g ((58%). Mp. 87°-90°C. Anal. for C₁₇H₇₅BrN₂O₇. Found (calc.): C: 42.33 (42.78); H: 5.15 (5.28); Br: 17.20 (16.74); N: 1.69 (11.74). H-NMR (DMSO-d, 250 MHz, 20 J in Hz): 1.93 & 11.92 (1H,s,C=ONHC=O); 8.09 & 8.07 (1H, s, C=C-H); 7.00 & 6.80 (1H, t, b, BocnH); 4.80 & 4.62 (2H,s,NCH,CON); 4.35 & 4.24 (2H,s,NCH,COOEt); 4.27-4.15 (2H, m's, COOCH, CH, O); 3.47-3.43 (2H, m's, BocNHCH, CH, N); 3.28-3.25 & 3.12-3.09 (2H, m's, BocNHCH, CH-, N): 1.46 € 25 (9H,s, Bu); 1.26 & 1.32 (3H,t,J=7.1, COOCH₂CH₃). 13C-NMR (DMSOd,, 250 MHz); 169.3 & 169.0 (EBuOC=0); 167.4 & 167.1 (COOEt); 159.8 (C=C-CON); 155.9 (NCH,CON); 150.4 (NCON); 145.9 (COCBr-CHN); 94.5 (COCBr=CHN); 78.2 (Me₁C); 61.3 & 60.7 (COCH₂CH₂); 49.1 & 48.0 (NCH,COOH); 48.0 & 47.0 (NCH,CON); 30 (BocNHCH, CH_2N); 38.2 (BocNH CH_2CH_2N); 26.3 (C(CH_3); _(COCH,CH₂). UV (Methanol; ___ NM): 226; 280. IR (KBr, CM⁻¹): 3200ms, broad (_NH); 168vs, vbroad (_C=0, COOH, CONH); 1250s (_ C-O, COOEt); 1170s (_C-O, COO Bu); 859m (d CH, Br-C=C-H). FAB-MS m/z (assignment, relative intensity): 479/477 (M + H, 35 5); 423/421 (M + 2H - Bu, 8); 379/377 (M + 2H - Boc, 100); 233/231 (M - backb ne, 20).

N-(B c-amin ethyl)-N-(5-br m uracyl-N¹-methylenecarbonoyl)-glycine

The product of Example 111 (1.96 g; 4.11 mmol) was 5 dissolved in methanol (30 ml) by heating, and then cooled t 0°C. Sodium hydroxide (2M, 30 ml) was added, and the mixture stirred for 30 min. HCl (1M, 70 ml) was added to pH = 2.0. The water phase was extracted with ethyl acetate (3 x 65 ml + 7 x 40 ml). The combined ethyl acetate extractions were 10 washed with saturated NaCl-solution (500 ml). acetate phase was dried over magnesium sulphate, filtered and evaporated to dryness in vacuo. Yield: 1.77 g (96%). Mp. 92 --97°C. Anal. for C₁₅H₂₁BrN₄O₇. Found (calc.): C: 40.79 (40.10); H: 5.15 (4.71); Br: 14.64 (17.70); N: 11.35 (12.47). 15 (DMSO-d_k, 250 MHz, J in Hz): 12.83 (1H,s,COOH); 11.93 & 11.91 (1H, s, C=ONHC=0); 8.10 & 8.07 (1H, s, C=C-H); 7.00 & 6.81 $(1H, t, b, Bocn_H)$; 4.79 & 4.61 $(2H, s, NC_{H_2}CON)$; 4.37 & 4.25 (2H,s,NCH2COOH); 3.46-3.39 (2H,m's, BocNHCH2CH2N); 3.26-3.23 £ 3.12-3.09 (2H, m's, BocNHCH, CH, N); 1.46 (9H, s, Bu). 15C-NMR 20 9DMSO-d₄,250 MHz); 170.4 (BuOC=0); 166.9(COOH); 159.7 (C=C-CON); 155.8 (NCH,CON); 150.4 (NCON); 145.9 (COCBr=CHN); 94.4 (COCBr=CHN); 78.1 (Me_C); 49.1 & 48.0 (NCH,COOH); 47.7 & 47.8 (NCH,CON); 38.6 (BOCNHC,CH,N); 38.1 (BOC NHCH,CH,N); 28.2 $(C(\underline{CH_3})_3)$. UV (Methanol; ___nm); 226; 278. IR (KBr,cm⁻¹): 25 3194ms, broad (_NH); 1686vs, vbroad (_C=O COOH, CONH); 1250s (_C-O,COOH); 1170s (_C-O,COO^TBu); 863m (& CH, Br-C=C-H). FAB-MS m/z (assignment, relative intensity): 449/451 (M + H, 70); 349/351 (M + 2H -Boc, 100); 231/233 (M - backbone, 20).

30 EXAMPLE 113

Uracil-N'-methyl acetate

Uracil (10.0 g; 89.2 mmol) and potassium carbonate (24.7 g; 178 mmol) were suspended in DMF (250 ml). Methyl bromoacetate (8.45 ml; 89.2 mmol) was added over a period f 35 5 min. The suspension was stirred overnight under nitr gen at ro m temperature, and then filt r d. TLC (10% methanol in ethylene chl ride) indicat d incomplete c nversion of uracil.

The solid residue was washed twice with DMF, and the combined filtrates were evaporated to dryness in vacuo. precipitate was suspended in water (60 ml) and HCl (2.5 ml, 4M) was added to pH = 2. The suspension was stirred for 30 5 min at 0°C, and then filtered. The precipitated title compound was washed with water and dried, in vacuo, sicapent. Yield: 9.91 g (60%). Mp. 1820 - 183°C. Anal. for $C_6H_8N_2O_4$. Found (calc.): C: 45.38 (45.66); H: 4.29 (4.38); N: 'H-NMR (DMSO-d, 250 MHz, J in Hz): 15.00 (15.21). 10 (1H,s, NH); 7.68 (1H,d, $J_{H-C=C-H}=7.9$), CH=CHN); 5.69 (1H,d, $J_{H-C=C-H}=7.9$) $_{H}$ =7.9), CH=CHN); 4.59 (2H,s,NCH₂COOMe); 3.76 (3H,s,COOCH₂). 13 C-NMR (DMSO-d₆, 250 MHz); 168.8 (COOMe); 164.0 (C=C-CON); 151.1 (NCON); 146.1 (COCH=CHN); 101.3 (COCH=CHN); 52.5 (COOCH₃); 48.7 (NCH₂COOMe). UV (Methanol; mrnm): 226; 261. 15 IR (KBr; cm⁻¹); 3164s (_NH); 1748vs (_C=0, COOMe); 1733vs (_C=0, CONH); 1450vs (∂ CH, CH₃0); 1243VS (_C-0,COOMe); 701m (∂ CH, H-C=C-H). FAB-MS m/z (assignment); 185 (M+H).

EXAMPLE 114

20 Uracilacetic acid

Water (90 ml) was added to the product of Example 113 (8.76 g; 47.5 mmol), followed by sodium hydroxide (2M, 40 ml). The mixture was heated for 40 min, until all the methyl ester has reacted. After stirring at 0°C for 15 min, hydrochloric 25 acid (4M, 25 ml) was added to pH=2. The title compound precipitated and the mixture was filtered after 2-3 h. precipitate was washed once with the mother liquor and twice with cold water and dried in vacuo over sicapent. Yield: Mp. 288°-289°C. Anal. for C,H,N,O,. 6,.66 g (82%). 30 (calc.): C: 42.10 (42.36), H: 3.43 (3.55); N: 16.25 (16.47)/ $^{1}H-NMR$ (DMSO- d_{A}), 250 MHz, J in Hz): 13.19 (1H,s,COOH); 11.41 (1H, s, NH); 7.69 $(1H, d, J_{H-C_{n}C_{n}H}=7.8, J_{H-C_{n}C_{n}H}=2.0, coch=chn)$; 4.49 (2H, s, NCH, COOH). 13C-NMR (DMSO-d₄, 2509 MHz); 169.9 (COOH); 163.9 (CH=CHCON); 151.1 (NCON); 146.1 (COCH=CHN); 100.9 35 (COCH=CHN); 48.7 NCH2COOH. UV (Methanol; ___nm): 246; 263. IR (KBr; cm⁻¹): 3122s (_NH); 1703vs (_C=0, COOH); 1698vs,

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1692vs (_C=0, CONH); 1205s (_C-0,COOH); 676 (d CH, H-C=C-H).

FAB-MS m/z (assignment): 171 (M + H).

EXAMPLE 115

5 N-(Bocaminoethyl)-N-(uracil-N'-methylenecarbonoyl)glycine ethyl ester

(Bocaminoethyl) glycine ethyl ester (2.00 g; 8.12 mmol) was dissolved in DMF (10 ml). Dhbt-OH (1.46 g; 8.93 mmol) was added and a precipitate was formed. DMF (2 x 10 ml) was added 10 until all was dissolved. The product of Example 114 (1.52 g; 8.93 mmol) was added slowly to avoid precipitation. Methylene chloride (30 ml) was added, and the mixture was cooled to 0°C, whereafter DDC (2.01g; 9.74 mmol) was added. The mixture was stirred for 1 h at 0°C, at 2 h at room temperature, and then The precipitated DCU was washed twice with 15 filtered. methylene chloride. To combined filtrate was added methylene chloride (100 ml), and the solution washed with half-saturat d NaHCO3-solution (3 x 100 ml, H2O:saturated NaHCO3-solution 1:1 v/v), then with dilute KHSO, solution (2 x 100 ml, 20 H₂O:saturated KHSO₄-solution 4:1 v/v) and finally with saturated NaCl-solution (1 x 100 ml). The organic phase was dried over magnesium sulphate, filtered and evaporated t dryness in vacuo (about 15 mmHg and then about 1mmHg). The residue was suspended in methylene chloride (32 ml), and 25 stirred for 35 min at room temperature, and 30 min at 0°C, and The precipitate (DCU) was washed with then filtered. methylene chloride. Petroleum ether (2 volumes) was added dropwise to the combined filtrate at 0°C, which caus d separation of an oil. The mixture was decanted, the remaining 30 oil was then dissolved in methylene chloride (20 ml), and th n again precipitated by addition of petroleum ether (2 volum s). This procedure was repeated 5 times until an impurity was removed. The impurity can be seen by TLC with 10% MeOH/CH,Cl, as the developing solvent. The resulting oil was dissolved 35 in methylene chloride (20 ml) and evaporated to dryness in vacuo, which caused solidificati n f the title c mpound. Yield: 1.71 g (53%). Mp. 68.5° - 75.7°C. Anal f r $C_{17}H_{24}N_{4}O_{7}$.

Found (calc.): C: 50.61 (51.25); H: 6.48 (6.58); N: 13.33 (14.06). $^{1}H-NMR$ (DMSO- d_{6} , 250 MHz, J in Hz): (1H,s,C=ONHC=0); 7.51 & 7.47 (1H,d, $J_{u-c=c-u}$ + 6.1; COCH=X-H); 7.00 & 6.80 (1H,t,b, BocNH); 5.83 & 5.66 (1H,d, $J_{u-c=c-x}=$ 5.7, 4.78 & 4.60 (2H,s,NCH,CON); 4.37 (2H,s,NCh,COOEt); 4.30 - 4.15 (2H,m's,COOCH,CH3); 3.49-3.46 (2H, m's, BocNHCH, CH, n); 3.27 3.23 & 3.11-3.09 (2H, m's, BOCNHCH,CH,N; 1.46 (9H, s, Bu); 1.39-1.23 (3H, m's, COOCH,CH,). ¹³C-NMR (DMSO-d₆, 250 MHz): 169.4 & 169.0 (¹BuOQ=0); 167.6 & 10 167.3 (COOEt); 163.8 (CH=CHCON); 155.8 (NCH,CON); 151.0 (NCON); 146.3 (COCH=CHN); 100.8 (COCH=CHN); 78.1 (Me.C); 61.2 & 60.6 (COOCH,CH3); 49.1 (NCH,COOEt); 47.8 & 47.0 (NCH,CON); 38.6 (BocNHCH₂CH₂N); 38.1 & 37.7 (BocNHCH₂N); 28.2 (C(CH₂)₃); 14.1 (CO-OCH₂CH₃. UV (Methanol; __ nm); 226; 264. IR (KBr; cm 15 1): 3053m (_NH); 1685vs, vbroad (_C=0, COOH, CONH); 1253s { C-0, COOEt); 1172s (_C-0, COO^TBu); 718w (d CH, C-C-C-H), FAB-MS m/z (assignment, relative intensity); 399 (M + H, 35); 343 (M + 2H - Bu, 100); 299 (M + 2H - Boc, 100); 153 (M-backbone, 30).

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EXAMPLE 116

N-(Bocaminoethyl)-N-(uracilmethylenecarbonoyl)glycine

The product of Example 115 (1.56 g; 3.91 mmol) was dissolved in methanol (20 ml) and then cooled to 0°C. Sodium 25 hydroxide (2M, 20 ml) was added, and the mixture was stirred for 75 min at 0°C. Hydrochloric acid (1M, 46 ml) was added to pH = 2.0. The water phase was extracted was ethyl acetate $(3 \times 50 \text{ ml} + 7 \times 30 \text{ ml}).$ The combined ethyl acetate extractions were washed with saturated NaCl solution (360 ml). 30 The ethyl acetate phase was dried over magnesium sulphate, filtered, and evaporated to dryness, in vacuo. The residue was dissolved in methanol and evaporated to dryness, in vacuo. Yield: 0.55 g (38%). Mp 164° - 170°C. Anal. for $C_{15}H_{22}N_{2}O_{7}$. Found (calc.): C: 46.68 (48.65); H: 6.03 (5.99); N: 1461 35 (15.13). H-NMR (DMSO-d, 250 MHz, J in Hz); 12.83 (1H, s, COOH); 11.36 (1H, s, C=ONHC=O); 7.52-7.45 (1H, m's, COCH=CHN); 7.00 & 6.82 (1H, t,b, BocNH); 5.67-5.62 (1H, M's, COCH=CHN);

4.76 & 4.58 (2H, s, NCH₂CON); 4.26 & 4.05 (2H, s, NCH₂COOH);
3.46-3.39 (2H, m's, BocNHCH₂CH₂N); 3.25-3.23 & 3.15-3.09 (2H, m's, BocNHCH₂CH₂N); 1.46 (9H, s, ^bBu). ¹³C-NMR (DMSO-d₆, 250 MHz); 170.5 (^bBuOg=O); 167.2 (COOH); 163.9 (C=C-QON); 155.8
5 (NCH₂CON); 151.1 (NgON); 146.4 (COCH=CHN); 100.8 (COCH=CHN); 78.1 (Me₂C); 49.1 & 47.8 (NgH₂ COOH); 47.6 & 46.9 (NgH₂CON); 38.6 (BocNHCH₂CH₂N); 38.1 & 37.6 (BocNHCH₂CH₂N); 28.2 (C(GH₂)₂). UV (Methanol; mm); 226; 264. IR (KBr; cm⁻¹); 3190 (_NH); 1685vs, vbroad (_C=O, COOH, CONH); 1253s (_C-O, COOH); 1171s (_C-O, COO^bBU); 682v (3 CH, H-C=C-H). FAB-NS m/z (assignm nt, relative intensity): 371 (M + H, 25); 271 (M + H -Boc, 100). EXAMPLE 117
H-U10-LysNH₂

Synthesis of the title compound was accomplished by using "synthetic Protocol 10". The synthesis was initiat d on approximately 100 mg Lys (Cl2)-MHBA-resin. The crude product (12 mg) was pure enough for hybridization studi s. The hybrid between 5'-(dA)10 and R-U10 had Tm of 67.5°C.

- 20 Deprotection and Cleavage of H-[Cacg]₁₀-Lys-NH₁ by Trifluoromethansulfonic Acid (TFMSA). An Alternative Meth d to Deprotection and Cleavage by Hydrogen Fluoride (HF).
 - (a) Deprotection of Side-Chain Protecting Groups
 by a "Low-Acidity" TFMSA-TFA-DMS Procedure
- A portion of ca. 0.4 g wet Boc-[Cacg]₁₀-Lys(CIZ)-MBHA resin (prepared in one of the previous examples) was pla ed in a 5 ml solid-phase reaction vessel. The n-Terminal Boc group was removed by the following protocol: (1) 50% TFA/CH₃Cl₂, 2 x 1 min and 1 x 30 min; (2) 100% TFA, 2 x 1 min and drain. In order to deprotect the benzyl-based side-chain protecting groups a so-called "low-acidity" TFMSA proc dure was carried out as follows: A stock solution (a) containing 5 ml of TFA-DMS-m-cresol (2:6:2, v/v/v) and a stock solution (B) containing TFA-TFMSA (8:2, v/v) were prepared. Next, the f llowing st ps were carried out: (3) 1 ml f stock solution (A) is add d to th PNA-resin in the rection vessel with shaking for 2 min. No drain; (4) 1 ml f stock solution (B) (cooled with ice/water) is add d in portions of 200 μl every

5

10th minute ver a p riod of 40 min, and shaking is c ntinu d for another 50 min; (5) drain and washing with 100% TFA, 5 x 1 min, and drain.

(b) Cleavage from the Resin by a "High-Acidity"
TPMSA-TFA Procedure

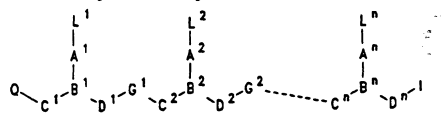
In order to cleave the above-mentioned deprotected PNA from the resin a so-called "high acidity" TFMSA procedure was carried out as follows: A stock solution (C) containing mcresol-TFA (2:8, v/v) was prepared. Next, the following steps 10 were carried out: (6) 1 ml of stock solution (C) was added t the deprotected PNA-resin in the SPPS vessel with shaking for 2 min; (7) 1 ml of stock solution (B) (cooled with ice/water) is added in portions of 200 μ l over a period of 30 min and shaking is continued for another 150 min; (5) the 2 ml solutin 15 in the reaction vessel is "blown out" through the filter int ml solution of diethylether cooled with dry ice/isopropanol. In order to complete the precipitati n process, 200 µl of anhydrous pyridine is added dropwise to the acid-ether mixture; (8) centrifugalization at 3000 rpm for 5 20 min; (9) the supernatant is decanted and the precipitat is washed three times with cold diethylether, dried, dissolv d in water, and lyophilized.

- (c) Purification and Identification of H[Caeg] 10-Lys-NH2
- An analytical HPLC chromatogram showed a nice crude product of good purity and a profile almost identical to that obtained from the HF cleavage of H-[Caeg]₁₀-Lys-NH₂, except that an additional peak, of course, arising from pyridine TFMSA salt elutes early in the chromatogram. Purificati n and identification was carried out by the usual procedures.

Those skilled in the art will appreciate that numer us changes and modifications may be made to the preferr d embodiments of the invention and that such changs and m dific tions may be made without departing from the spirit of the inv nti n. It is ther fore intended that the appended claims cov r all such equival nt variations as fall within the tru spirit and so pe f the inventi n.

WHAT IS CLAIMED IS:

- 1. A ompound c mprising a polyamide backb ne bearing a plurality of ligands that are individually bound to aza nitrogen atoms located within said backbone, at least on f said ligands being a naturally occurring nucleobase, a nnnaturally occurring nucleobase, a DNA intercalator, r a nucleobase-binding group.
- 2. The compound of claim 1 wherein said aza nitr gen atoms are separated from one another in said backbone by from 4 to 6 intervening atoms.
 - 3. A compound having the formula:

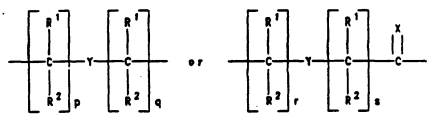


wherein:

n is at least 2,

each of L^1-L^n is independently selected from the gr up consisting of hydrogen, hydroxy, (C_1-C_4) alkanoyl, naturally occurring nucleobases, non-naturally occurring nucleobases, aromatic moieties, DNA intercalators, nucleobase-binding groups, heterocyclic moieties, and reporter ligands, at 1 ast one of L^1-L^n being a naturally occurring nucleobase, a non-naturally occurring nucleobase, a DNA intercalator, r a nucleobase-binding group;

each of A^1-A^n is a single bond, a methylene group r a group of formula:



where:

X is O, S, Se, NR³, CH₂ or C(CH₃)₂; Y is a single bond, O, S or NR⁴;

each of p and q is zero or an integer from 1 to 5, the sum p+q being not more than 10;

each of r and s is zero or an integer from 1 to 5, the sum r+s being not more than 10;

each R^1 and R^2 is independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl which may be hydroxy- or alkoxy- or alkylthio-substituted, hydroxy, alkylthio, amino and halogen; and

each R^3 and R^4 is independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl, hydroxy- or alkoxy- or alkylthio-substituted (C_1-C_4) alkyl, hydroxy, alkoxy, alkylthio and amino;

each of B^1-B^n is N or R^3N^4 , where R^3 is as defined above;

each of C^1-C^n is CR^6R^7 , CHR^6CHR^7 or $CR^6R^7CH_2$, where R^6 is hydrogen and R^7 is selected from the group consisting of the side chains of naturally occurring alpha amino acids, or R^6 and R^7 are independently selected from the group consisting of hydrogen, (C_2-C_6) alkyl, aryl, aralkyl, heteroaryl, hydroxy, (C_1-C_6) alkoxy, (C_1-C_6) alkylthio, NR^3R^6 and SR^5 , where R^3 and R^6 are as defined above, and R^5 is hydrogen, (C_1-C_6) alkyl, hydroxy-, alkoxy-, or alkylthio- substituted (C_1-C_6) alkyl, R^6 and R^7 taken together complete an alicyclic or heterocyclic system;

each of D^1-D^n is CR^6R^7 , $CH_2CR^6R^7$ or CHR^6CHR^7 , where R^6 and R^7 are as defined above;

each of G^1-G^{n-1} is $-NR^3CO-$, $-NR^3CS-$, $-NR^3SO-$ or $-NR^3SO_2-$, in either orientation, where R^3 is as defined above;

Q is $-CO_2H$, -CONR'R'', $-SO_3H$ or $-SO_2NR'R''$ or an activated derivative of $-CO_2H$ or $-SO_3H$; and

I is -NHR'''R'''' or -NR'''C(0)R'''', where R', R", R''' and R'''' are independently selected from the group c nsisting f hydrogen, alkyl, amino protecting gr ups, reporter ligands, intercalators, chelators, peptides,

proteins, carbohydrates, lipids, steroids, ligonucl otides and soluble and non-solubl polymers.

4. The compound of claim 3 having the formula:

wherein:

each L is independently selected from the group consisting of hydrogen, phenyl, heterocyclic moi ti s, naturally occurring nucleobases, and non-naturally occurring nucleobases;

each $R^{7'}$ is independently selected from the group consisting of hydrogen and the side chains of naturally occurring alpha amino acids;

n is an integer from 1 to 60, each k and m is, independently, zero or 1; each 1 is zero or an integer from 1 to 5; Rh is OH, NH₂ or -NHLysNH₂; and Ri is H or COCH₃.

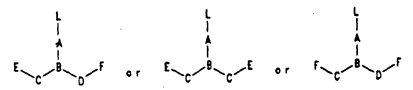
5. The compound of claim 4 having formula:

-148-

each L is independently selected from the group consisting of the nucleobases thymine, adenine, cytosine, guanine, and uracil;

each R' is hydrogen; and n is an integer from 1 to 30.

6. A compound having one of the following formulas:



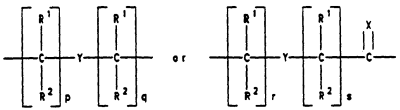
wherein:

L is selected from the group consisting of hydrogen, hydroxy, (C_1-C_ℓ) alkanoyl, naturally occurring nucleobases, non-naturally occurring nucleobases, aromatic moieties, DNA intercalators, nucleobase-binding groups, and heterocyclic moieties, reporter ligands, wherein:

at least one of L¹-Lⁿ is a naturally occurring nucleobase, a non-naturally occurring nucleobase, a DNA intercalator, or a nucleobase-binding group; and

amino groups are, optionally, protected by amino protecting groups;

A is a single bond or a group of the formula:



where:

X is O, S, Se, NR³, CH₂ or C(CH₃)₂; Y is a single bond, O, S or NR⁴;

each of p and q is zero or an integer from 1 to
5, the sum p+q being not more than 10;

each f r and s is zero r an int ger fr m 1 to 5, the sum r+s being not mor than 10;

each R^1 and R^2 is independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl, hydroxy- ralkoxy- or alkylthio-substituted (C_1-C_4) alkyl, hydroxy, alkylthio, amino and halogen; and

each R^3 and R^4 is independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl, hydroxy-ralkoxy-or alkylthio-substituted (C_1-C_4) alkyl, hydroxy, alkylthio and amino;

B is N or R^3N^4 , where R^3 is as defined above;

each C is CR^6R^7 , CHR^6CHR^7 or $CR^6R^7CH_2$, where R^6 is hydrogen and R^7 is selected from the group consisting of the side chains of naturally occurring alpha amino acids, or R^6 and R^7 are independently selected from the group consisting of hydrogen, (C_2-C_6) alkyl, aryl, aralkyl, heteroaryl, hydr xy, (C_1-C_6) alkoxy, (C_1-C_6) alkylthio, NR^3R^6 and SR^5 , where R^3 and R^6 are as defined above, and R^5 is hydrogen or (C_1-C_6) alkyl, hydroxy-, alkoxy-, or alkylthio- substituted (C_1-C_6) alkyl, R^6 and R^7 taken together complete an alicyclic or heterocyclic system;

each D is CR^6R^7 , $CH_2CR^6R^7$ or CHR^6CHR^7 , where R^6 and R^7 are as defined above;

each E is COOH, CSOH, SOOH, SO₂OH or an activated r protected derivative thereof; and

each F is NHR³ or NPgR³, where R³ is as defined above, and Pg is an amino protecting group.

7. The compound of claim 6 having the formula:

wherein:

each L is independently selected from the group consisting of hydrogen, phenyl, heterocyclic moieties, naturally occurring nucleobases, and non-naturally occurring nucleobases;

each $R^{7\prime}$ is independently selected from the group consisting of hydrogen and the side chains of naturally occurring alpha amino acids; and

each k, 1, and m is, independently, zero or an integer from 1 to 5.

8. The compound of claim 7 having formula:

wherein:

L is selected from the group consisting of the nucleobases thymine, adenine, cytosine, guanine, uracil, 5-methylcytosine, 6-thioguanine and 5-bromouracil, and prot cted derivatives thereof;

R' is hydr g n;

E is COOH or an activated or pr tected derivative thereof; and

F is NH_2 or NHPg , where Pg is an amino protecting group.

9. A compound having the formula:

wherein:

each L is independently selected from the group consisting of hydrogen, phenyl, heterocyclic moieties, naturally occurring nucleobases, and non-naturally occurring nucleobases;

each $R^{7'}$ is independently selected from the grup consisting of hydrogen and the side chains of naturally occurring alpha amino acids;

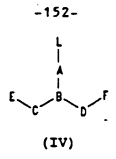
n is an integer from 1 to 60,

each k, 1, and m is, independently, zero or an int ger
from 1 to 5;

 R^h is OH, NH_2 or $-NHLysNH_2$; and

Ri is H or COCH,

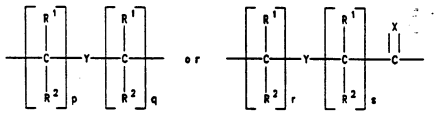
- 10. A process for preparing a compound according to claim 1, comprising the steps of:
- A) providing a polymer substrate, said polymer being functionalized with a chemical group capable of forming an anchoring linkage with an amino acid;
- B) coupling said polymer with a first amino acid through said anchoring linkag, said first amino acid having formula (IV):



wherein:

L is selected from the group consisting of naturally occurring nucleobases, non-naturally occurring nucleobases, aromatic moieties, DNA intercalators, nucleobase-binding groups, heterocyclic moieties, and reporter ligands, wherein amino groups are, optionally, protected by amino protecting groups;

A is a single bond or a group of the formula:



where:

X is 0, S, Se, NR^3 , CH_2 or $C(CH_3)_2$;

Y is a single bond, O, S or NR⁴;

p and q are zero or integers from 1 to 5, the sum
p+q being not more than 10;

r and s are zero or integers from 1 to 5, the sum r+s being not more than 10;

 R^1 and R^2 are independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl, hydroxy- or alkoxy- or alkylthio-substituted (C_1-C_4) alkyl, hydroxy, alkoxy, alkylthio, amino and halogen; and

 R^3 and R^4 are independently selected from the group consisting of hydrogen, (C_1-C_4) alkyl, hydroxy- or alkoxy- or alkylthio-substituted (C_1-C_4) alkyl, hydroxy, alkoxy, alkylthio and amino;

B is N r R³N^{*}, wh r R³ is as defined ab ve;

C is CR^6R^7 , CHR^6CHR^7 r $CR^6R^7CH_2$, where R^6 is hydrogen and R^7 is selected fr m th group consisting

f the side chains of naturally courring alpha amino acids, or R^6 and R^7 are ind p ndently selected from the group consisting of hydrogen, (C_2-C_6) alkyl, aryl, aralkyl, heteroaryl, hydroxy, (C_1-C_6) alkoxy, (C_1-C_6) alkylthio, NR^3R^4 and SR^5 , where R^3 and R^4 are as defined above, and R^5 is hydrogen or (C_1-C_6) alkyl, hydroxy-, alkoxy-, or alkylthio- substituted (C_1-C_6) alkyl, or R^6 and R^7 taken together complet an alicyclic or heterocyclic system;

D is CR⁶R⁷, CH₂CR⁶R⁷ or CHR⁶CHR⁷, where R⁶ and R⁷ are as defined above;

E is COOH or an activated or protected derivative thereof; and

F is NPgR³ where R³ is as defined above and Pg is an amino protecting group;

- C) removing said amino protecting group from said coupled first amino acid to generate a free amino group; and
- D) reacting said free amino group with a second amino acid having formula (IV) to form a peptide chain.
- 11. The process of claim 10 further comprising the steps of:
- E) removing said amino protecting group from said second amino acid to generate a terminal free amino group on said peptide chain; and
- F) reacting said free amino group on said peptide chain with a further amino acid having formula (IV) to lengthen said peptide chain.
- 12. The process of claim 11 wherein steps E and F are performed a plurality of times.
 - 13. The process of claim 11 further comprising removing at least one protecting group remaining on the amin acid moieties of the peptide chain.

- 14. The process of claim 10 further comprising cleaving said anchoring linkage without substantially degrading said peptide chain.
- 15. The process of claim 10 wherein the polymer substrate contains polystyrene, polyacrylamide, silica, a composite material, cotton, or a derivative thereof.
- 16. The process of claim 10 wherein the chemical group capable of forming said anchoring linkage is chloro-, bromo- and iodo-substituted alkyl, amino-substituted alkyl, amino and aryl-substituted alkyl, amino- and alkylaryl-substituted alkyl, hydroxy-substituted alkyl, or a derivative thereof having a spacer group that can be cleaved substantially without degradation of said polypeptide.
- 17. The process of claim 16 wherein chloro-substituted alkyl is chloromethyl, amino-substituted alkyl is aminomethyl, amino- and alkyl-substituted aryl is α -aminobenzyl, amino- and alkylaryl-substituted alkyl is selected from the group consisting of α -amino-3- and α -amin-4-methylbenzyl, and hydroxy-substituted alkyl is hydroxymethyl.

18. The process of claim 16 wherein:

the chemical group is derived from an amino-containing moiety selected from amino-substituted alkyl, amino- and aryl substituted alkyl, and amino- and alkylaryl-substituted alkyl; and

the chemical group includes a spacer group derived from the group consisting of 4-(haloalkyl)aryl-lower alkanoic acids, Boc-aminoacyl-4-(oxymethyl)aryl-lower alkanoic acids, N-Boc-p-acylbenzhydrylamines, N-Boc-4'-(lower alkyl)-p-acylbenzhydrylamines, N-Boc-4'-(lower alkoxy)-p-acylbenzhydrylamines, and 4-hydroxymethylphenoxy-lower alkanoic acids.

- 19. A pr c ss f r s quence-sp cific rec gnition f a doubl -stranded polynucleotide, comprising c ntacting said polynucleotide with a compound that is diff r nt from natural RNA and that binds to one strand of the polynucleotide, thereby displacing the other strand.
- 20. The process of claim 19 wherein said compound is an oligomer comprising a homogenous or heterogenous backbone to which are linked naturally occurring nucleobases, n n-naturally occurring nucleobases or other ligands that individually bind by hydrogen to at least one natural nucleobase in said bound polynucleotide strand.
- 21. The process of claim 20 wherein said compound is the compound of claim 1.
- 22. The process of claim 20 wherein said compound is the compound of claim 4.
 - 23. A process for modulating the expression of a gen in an organism, comprising administering to said organism a compound according to claim 1 that specifically binds t DNA or RNA deriving from said gene.
 - 24. The process of claim 23 wherein said compound is the compound of claim 1.
 - 25. The process of claim 23 wherein said compound is the compound of claim 4.
 - 26. The process of claim 23 wherein said modulation includes inhibiting transcription of said gene.
 - 27. The process of claim 23 wherein said modulation includes inhibiting replication of said gene.

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- 28. A process for treating c nditions associated with undesired protein production in an organism, comprising contacting said organism with an effective amount of a compound according to claim 1 that specifically binds with DNA or RNA deriving from a gene controlling said protein production.
- 29. The process of claim 28 wherein said compound is the compound of claim 1.
- 30. The process of claim 28 wherein said compound is the compound of claim 4.
- 31. A process for inducing degradation of DNA or RNA in cells of an organism, comprising administering to said organism a compound according to claim 1 that specifically binds to said DNA or RNA.
- 32. A process for killing cells or virus, comprising contacting said cells or virus with a compound according to claim 1 that specifically binds to a portion of the genome of said cells or virus.
- 33. A pharmaceutical composition comprising a compound according to claim 1 and at least one pharmaceutically effective carrier, binder, thickener, diluent, buffer, preservative, or surface active agent.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/EP 92/01219

Box	1	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
Thi	inter	national search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.		Claims Nos.: because they reluz to subject matter not required to be searched by this Authority, namely: Remark: Although claims 23-31 are directed to a method of treatment of the human/animal body the search has been carried out and based on the alleged effects of the compounds.
2		Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	لــا	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Bo	x II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
73	is Int	ernational Searching Authority found multiple investions in this international application, as follows:
1.		As all required additional search fees were timely paid by the applicant, this international search report covers all searchable ciaims.
2	. [As all searchable claims could be searches without effert justifying an additional fee, this Authority did not invite payment of any additional fee.
,	. [As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
	۰. [No required additional search feet were timely public by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the dalant, it is covered by claims Not.:
	R.	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

EP 9201219 SA 60822

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 24/09/92

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Patent family member(s)		Pablication date	
WO-A- 8605518	25-09-86	AU-A- AU-A- CA-A- EP-A- EP-A- JP-T- JP-T- WO-A- US-A-	5661386 5698186 1268404 0216860 0215942 62502338 62502357 8605519 5034506	13-10-86 13-10-86 01-05-90 08-04-87 01-04-87 10-09-87 10-09-87 25-09-86 23-07-91	
WO-A- 8605519	25-09-86	AU-A- AU-A- CA-A- EP-A- EP-A- JP-T- JP-T- WO-A- US-A-	5661386 5698186 1268404 0216860 0215942 62502338 62502357 8605518 5034506	13-10-86 13-10-86 01-05-90 08-04-87 01-04-87 10-09-87 10-09-87 25-09-86 23-07-91	

INTERNATIONAL SEARCH REPORT

International Application No.

PCT/EP 92/01219

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III. DOCUM		D TO BE RELEVANT				
Category •	Citation of D	ocument, 11 with indication, where approp	rists, of the relevant passages 12	Relevant to Claim No		
X	Septem	605518 (J. SUMMERTON) ber 1986 (cited in the	e application)			
Y	Septem	8605519 (J. SUMMERTON) aber 1986				
P,X	Journal of the American Chemical Society, vol. 114, no. 5, 26 February 1992, American Chemical Society, M. EGHOLM et al.: "Peptide nucleic acids (PNA). Oligonucleotide analogues with an achiral peptide backbone", pages 1895-1897					
Ρ,Χ	et al stran	ce, vol. 254, 6 December: "Sequence-selective displacement with a mide", pages 1497-1500	recognition of DNA by thymine-substituted			
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-	IFICATION			- I Smark Bread		
		of the International Search	Date of Mailing of this Internation	THE DESIGNATION IN COLUMN TWO IS NOT THE OWNER.		

Purm PCT/ISA/210 (second about) (Jamery 1985)

Page 2 PCT/EP 92/01219

International Application No.

	International Application No PCT	/EP 92/01219 4
	TS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	
Crietosà .	Citation of Document, with Indication, where appropriate, of the relevant passages	Relevant to Claim No.
A	Journal of The American Chemical Society, vol. 110, no. 22, 26 October 1988, American Chemical Society, D.P. MACK et al.: "Design and chemical synthesis of a sequence-specific DNA-cleaving protein", pages 7572-7574	
A	Journal of Medicinal Chemistry, vol. 33, no. 7, 1990, American Chemical Society, L.P.G. WAKELIN et al.: "Kinetic and equilibrium binding studies of amsacrine-4-carboxamides: a class of asymmetrical DNA-intercalating agents which bind by threading through the DNA helix", pages 2039-2044	
A	BIOCHEMISTRY, vol. 27, no. 17, 23 August 1988, American Chemical Society, P.E. NIELSEN et al.: "Photochemical cleavage of DNA by nitrobenzamides linked to 9-aminoacridine", pages 6338-6343	
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COMPOUND

COMPOUND 1 IN 50% TFA: 50% METHYLENE CHLRIDE, 5 h, rt.

COMPOUND 1 IN 100% HF, 0°C, 1h

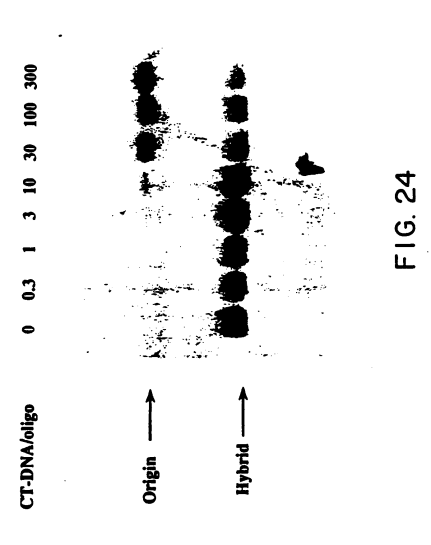
QUANTITATIVE DE-BENZYLATION

QUANTITATIVE DE-BENZYLATION AND DE-SULFONYLATION

Fig. 27

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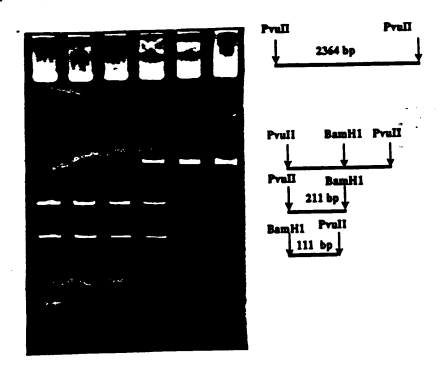
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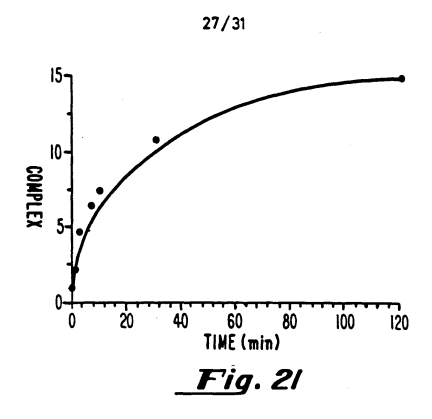
PNA/DNA 0 0.006 0.02 0.06 0.2 0.6

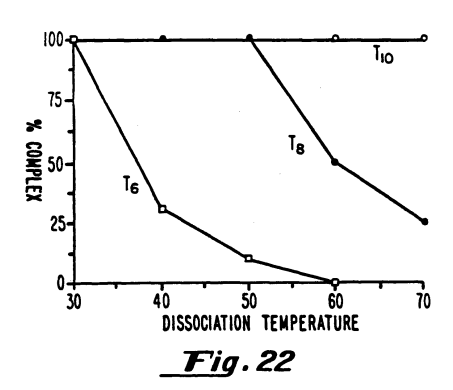


PNA Target

5'----GGATCCAAAAAAAAGGATCC----3'----CCTAGGTTTTTTTTTCCTAGG----BamH1
BamH1

FIG. 23

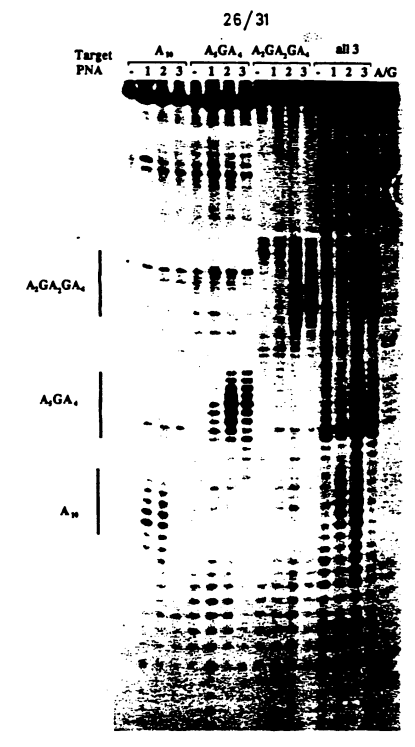




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PNA 1: T₁₀ PNA 2: T₅CT₄ PNA 3: T₂CT₅CT₄

FIG. 20
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Fig. 19

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Fig. 15

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Fig. 13

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S₁-nuclease 0.1 1 10 0.1 1 10 AcrT10Lys

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FIG. 12C

ACTTIOLYS - + ++ - + ++ - + + 3000m **HOO - steph 4/0 16/31

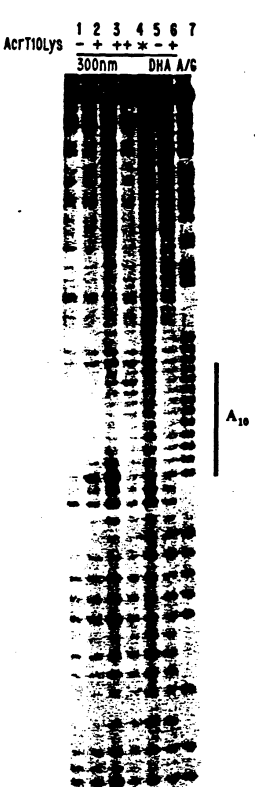


FIG. 12 B

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FIG. 12A

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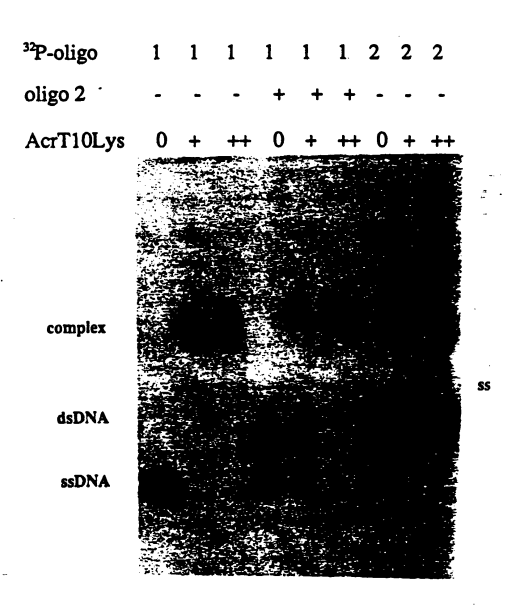


FIG. IIA

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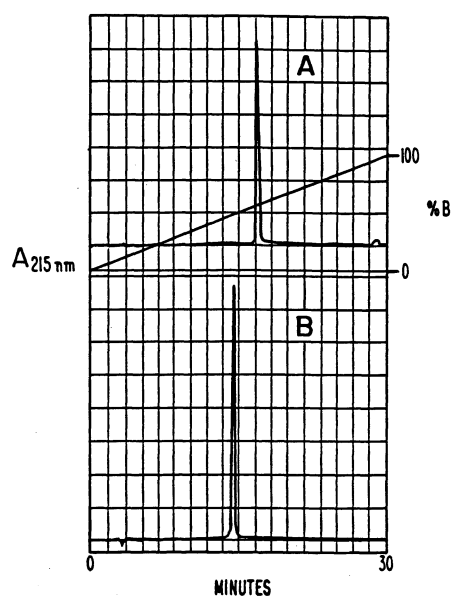


Fig. 10

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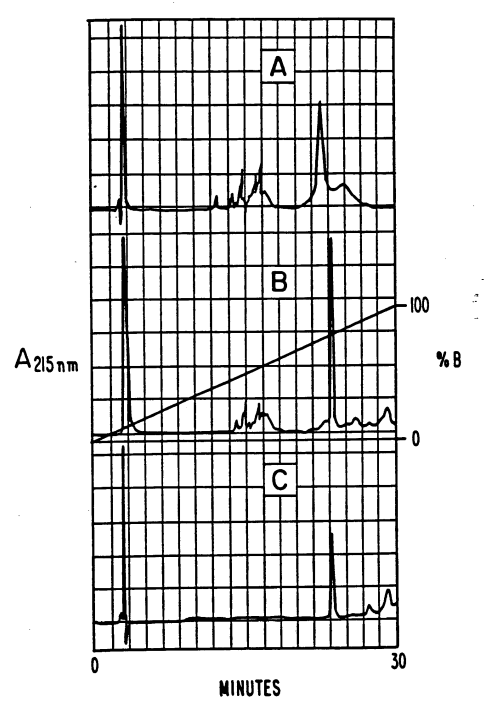
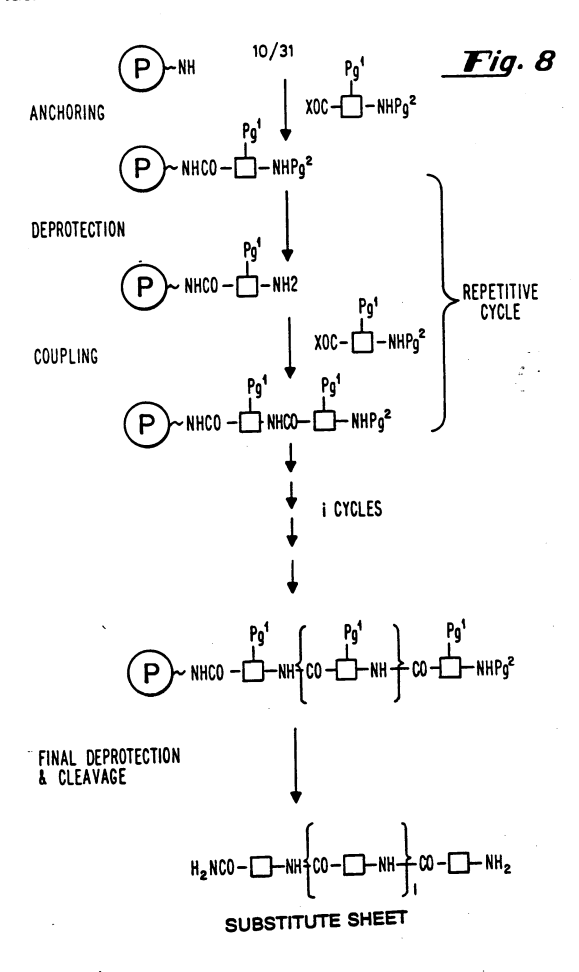


Fig. 9

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HN

NH-C

NO2

Acr
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<u>Fig. 5</u>

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Fig. 4

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KMnO4 CLEAVAGE

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Fig. 1(A)